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                 Source of Registration (SR) information in REGISTRY updated
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                 and searchable
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         JAN 27
                 A new search aid, the Company Name Thesaurus, available in
                 CA/CAplus
NEWS 5 FEB 05
                 German (DE) application and patent publication number format
                 changes
        MAR 03
NEWS
     6
                 MEDLINE and LMEDLINE reloaded
NEWS
     7
         MAR 03
                 MEDLINE file segment of TOXCENTER reloaded
NEWS
     8
         MAR 03
                FRANCEPAT now available on STN
NEWS
     9
         MAR 29
                 Pharmaceutical Substances (PS) now available on STN
NEWS 10
         MAR 29
                 WPIFV now available on STN
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         MAR 29
                 No connect hour charges in WPIFV until May 1, 2004
NEWS 12
         MAR 29
                 New monthly current-awareness alert (SDI) frequency in RAPRA
NEWS EXPRESS MARCH 31 CURRENT WINDOWS VERSION IS V7.00A, CURRENT
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 3 MARCH 2004
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             CAS World Wide Web Site (general information)
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FILE 'HOME' ENTERED AT 11:20:49 ON 13 APR 2004

=> file medline COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE LAST UPDATED: 10 APR 2004 (20040410/UP). FILE COVERS 1951 TO DATE.

On February 29, 2004, the 2004 MeSH terms were loaded. See HELP RLOAD for details. OLDMEDLINE now back to 1951.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2004 vocabulary. See http://www.nlm.nih.gov/mesh/ and http://www.nlm.nih.gov/pubs/techbull/nd03/nd03_mesh.html for a description of changes.

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
=> e cholinesterase
                      CHOLINESTERAS/BI
 E2
                      CHOLINESTERASA/BI
           18588 --> CHOLINESTERASE/BI
 E3
 E4
              1
                    CHOLINESTERASE2/BI
E5
                    CHOLINESTERASEACTIVITAS/BI
               1
              1 CHOLINESTERASEAKTIVITAET/BI
1 CHOLINESTERASEAKTIVITASA/BI
1 CHOLINESTERASEAKTIVITASANAK/BI
23 CHOLINESTERASEAKTIVITAT/BI
2 CHOLINESTERASEAKTIVITATEN/BI
E6
E7
E8
E9
E10
E11
                    CHOLINESTERASEAKTTVITAT/BI
               1
E12
               1
                      CHOLINESTERASEANTAGONISTEN/BI
=> s e1-e3
               4 CHOLINESTERAS/BI
               7 CHOLINESTERASA/BI
           18588 CHOLINESTERASE/BI
L1
          18590 (CHOLINESTERAS/BI OR CHOLINESTERASA/BI OR CHOLINESTERASE/BI)
=> e urinary
E1
               2
                      URINARUM/BI
E2
               2
                     URINARV/BI
E3
         200023 --> URINARY/BI
                  URINARYAD7C/BI
E4
              1
E5
               1
                     URINARYEXCRETION/BI
                    URINARYL/BI
E6
               1
                    URINARYNI/BI
E7
               1
E8
              1
                     URINARYS/BI
E9
               1
                     URINARYTOXICITY/BI
E10
               2
                     URINARYTRACT/BI
E11
               1
                     URINARYU/BI
E12
               1
                     URINARYVOLUME/BI
=> s e2 or e3 or e5 or e12
               2 URINARV/BI
         200023 URINARY/BI
               1 URINARYEXCRETION/BI
               1 URINARYVOLUME/BI
L_2
         200026 URINARV/BI OR URINARY/BI OR URINARYEXCRETION/BI OR URINARYVOLUME
                 /BI
=> s l1 and l2
            172 L1 AND L2
=> e muscarinic
E1
                     MUSCARINES/BI
E2
                     MUSCARINI/BI
E3
          21667 --> MUSCARINIC/BI
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09/ 960,477
            8 MUSCARINIC1/BI
3 MUSCARINIC2/BI
1 MUSCARINIC4/BI
2 MUSCARINICA/BI
20 MUSCARINICALLY/BI
1 MUSCARINICE/BI
3 MUSCARINICI/BI
3 MUSCARINICO/BI
 E4
 E5
 E6
 E7
 E8
 E9
 E10
 E11
 E12
              8
                     MUSCARINICOS/BI
 => s e3-e6
           21667 MUSCARINIC/BI
               8 MUSCARINIC1/BI
               3 MUSCARINIC2/BI
               1 MUSCARINIC4/BI
           21668 (MUSCARINIC/BI OR MUSCARINIC1/BI OR MUSCARINIC2/BI OR MUSCARINIC
L4
                  4/BI)
=> s 11 and 12 and 14
              13 L1 AND L2 AND L4
=> s (11 or 14) and 12
             712 (L1 OR L4) AND L2
=> s 16 and (dysuria or (urinary bladder))
            1700 DYSURIA
         200023 URINARY
          93807 BLADDER
          14036 URINARY BLADDER
                    (URINARY (W) BLADDER)
             327 L6 AND (DYSURIA OR (URINARY BLADDER))
L7
=> s 17 and inhibit?
        1095051 INHIBIT?
L8
             154 L7 AND INHIBIT?
=> s (l1 or l4) and inhibit?
        1095051 INHIBIT?
L9
          21065 (L1 OR L4) AND INHIBIT?
=> s 19 and 12
L10
            298 L9 AND L2
=> s 110 and (dysuria or bladder?)
           1700 DYSURIA
          94128 BLADDER?
L11
            198 L10 AND (DYSURIA OR BLADDER?)
=> s l11 not py>1998
       2690081 PY>1998
L12
            121 L11 NOT PY>1998
=> d l12 1- ibib abs
YOU HAVE REQUESTED DATA FROM 121 ANSWERS - CONTINUE? Y/(N):y
```

L12 ANSWER 1 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

AUTHOR CORPORATE SOURCE:

1 MEDLINE on STN
1999032785 MEDLINE
PUDMed ID: 9791087
M2 muscarinic receptor contributes to contraction
of the denervated rat urinary bladder.
Braverman A S; Luthin G R; Ruggieri M R
Department of Urology, Temple University School of
Medicine, Philadelphia, Pennsylvania 19140, USA.
RO1-DK-39086 (NIDDK) CONTRACT NUMBER

RO1-DK-43333 (NIDDK)

SOURCE:

American journal of physiology, (1998 Nov) 275 (5 Pt 2) R1654-60.

R1654-60. Journal code: 0370511. ISSN: 0002-9513. United States Journal; Article; (JOURNAL ARTICLE) English Priority Journals 199812 PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY MONTH:

ENTRY DATE:

E SEGMENT: Priority Journals RY MONTH: 199812 RY MONTH: 199812 Last Updated on STN: 1999015 Entered STN: 1999015 In vitro bladder contractions in response to cumulative carbachol doses were measured in the presence of selective muscarinic antagonists from rats that had their major pelvic gaugalion bilaterally removed. Denervation induced both hypertrophy and a supersensitivity of the bladders to agonist. The affinities in control bladders for antagonism of carbachol-induced contractions were consistent with M3-mediated contractions. Affinities in denervated bladders for 4-diphenlacetoxy-N methylpiperidine methiodide (8.5) and p-fluoro hexahydrosilodifenidol (6.6) were consistent with M3-mediated contractions. Subtype-selective immunoprecipitation of muscarinic receptors revealed a 50% increase in total and a 60% increase in the Y receptor density with no change in M3 receptor density in denervated bladders compared with normal or sham-operated controls. This increase in M2 receptor density is consistent with the change in affinity of the antagonists for inhibition of carbachol-induced contractions and may indicate that M2 receptors or a combination of M2 and M3 receptors directly mediates smooth muscle contraction in the denervated bladder.

ANSWER 3 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

1 MEDLING ON SIN 1998405233 MEDLINE PubMed ID: 9734316 Selective muscarinic antagonists. II. Synthesis and antimuscarinic properties of biphenylylcarbamate

AUTHOR:

and antimuscarinic properties of biphenylylcarbamate derivatives.
Naito R; Takeuchi M; Morihira K; Hayakawa M; Ikeda K; Shibanuma T; Isomura Y
Institute for Drug Discovery Research, Yamanouchi Pharmaceutical Co., Ltd., Ibaraki, Japan.
Chemical & pharmaceutical bulletin, (1998 Aug) 46 (8) 1286-94.

CORPORATE SOURCE:

SOURCE:

Journal code: 0377775, ISSN: 0009-2363. PUB. COUNTRY:

Journal code.

Japan

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE: LANGUAGE:

English Priority Journals 199810 FILE SEGMENT: ENTRY MONTH:

E SECRET: Priority Journals

RY MONTH: 199810

RY MONTH: 199810

Last Updated on STN: 19981020

Entered STN: 19981020

Last Updated on STN: 19981020

Entered Medline: 19981020

A novel series of biphenylylcarbamate derivatives were synthesized and evaluated for binding to Mi. M2 and M3 receptors and for antimuscarinic activities. Receptor binding assays indicated that biphenyl-2-ylcarbamate derivatives had high affinities for M1 and M3 receptors and good selectivities for M3 receptor over W2 receptor, indicating that the biphenyl-2-yl group is a novel hydrophobic replacement for the benzhydryl group in the muscarinic antagonist field. In this series, quinuculdini-4-yl biphenyl-2-ylcarbamate monohydrochloride (sl. YM-46303) exhibited the highest affinities for M1 and M3 receptors, and selectivity for M3 over M2 receptor. Compared to oxybutynin, YM-46303 showed approximately ten times higher inhibitory activity on bladder pressure in reflexly-evoked rhythmic contraction, and about 5 fold greater selectivity for winary bladder contraction against salivary secretion in rats. Moreover, selective antagonistic activity was also observed in vitro. Further evaluation of antimuscarinic effects on bradycardia and pressor in pithed rats, and on tremor in mice, showed that YM-46303 can be useful for the treatment of urinary urge incontinence as a bladder-selective M3 antagonist with potent activities and fewer side effects.

L12 ANSWER 2 OF 121 ACCESSION NUMBER: 1 DOCUMENT NUMBER: 1

1 MEDLINE on STN
1998420502 MEDLINE
PubMed ID: 9748713
Comparison of the effects of various spasmolytic drugs on isolated human and porcine detrusor smooth muscle.
Uckert S; Stief C G; Odenthal K P; Becker A J; Truss M C; Jonas U

AUTHOR:

Jonas U

Hannover Medical School, Department of Urology, Germany,
Arzneimittel-Porschung, (1998 Aug) 48 (8) 836 9.
JOurnal code: 0372660. ISSN: 0004-4172.
GERMANY: Germany, Federal Republic of
Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals
199811
Entered STM. -CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

ENTRY DATE:

ESSCHENT: Priority Journals
RY MONTH: 199311
RY MONTH: 199311
RY DATE: Entered STN: 19990106
Last Updated on STN: 19990106
Entered Medline: 19981109
The spasmolytic activity of flavoxate (CAS 15301-69-6), anticholinergic agents oxybutymin (CAS 56;33-20-5), and trospium chloride (CAS 10405-02-4), drugs commonly utilized in the therapy of hyperactive bladder, and phosphodesterase (PDE) inhibitors papaverine (CAS 58-74-2) and vinpocetine (CAS 42971-09-5) on muscarinic contractions of detrusor smooth muscle strips isolated from human and porcine urinary bladder was studied in vitro using the organ bath technique. Trospium chloride was most effective in relaxing contractions elicited by muscarinic stimulation, while flavoxate was significantly less effective than all other drugs tested. The relaxing potency of oxybutymin was greater than those of PDE-inhibitors papaverine and vinpocetine but 3,000 fold less significant than those of trospium chloride. The effects of the individual drugs on muscarinic tension of both human and porcine detrusor muscle strips were nearly equal. The present results suggest that the pig might be an appropriate animal model for the study of effects of spasmolytic substances on the contractility of urinary bladder smooth muscle in vitro.

L12 ANSWER 4 OF 121 ACCESSION NUMBER: MEDLINE on STN 1998405232 MEDLINE PubMed ID: 9734315 DOCUMENT NUMBER:

TITLE:

PubMed ID: 9734315 Selective muscarinic antagonists. I. Synthesis and antimuscarinic properties of 4-piperidyl benzhydrylcarbamate derivatives. Naito R, Takeuchi M, Morihira K; Hayakawa M; Ikeda K; Shibanuma T; Isomura Y Institute for Drug Discovery Research, Yamanouchi Pharmaceutical Co., Ltd., Ibaraki, Japan. Chemical & pharmaceutical bulletin, (1998 Aug) 46 (8) 1274-85. AUTHOR:

CORPORATE SOURCE:

SOURCE:

Journal code: 0377775. ISSN: 0009-2363

Journal code: 0377775. ISSN: 0009-2 Japan Journal; Article; (JOURNAL ARTICLE) English Priority Journals 193810 PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT:

ENTRY MONTH

ENTRY DATE

SEGMENT: Priority Journals

Y MONTH: 199810

Entered STN: 19981020

Last Updated on STN: 19981020

Entered Meddine: 19981002

A series of 1-substituted-4-piperidyl benzhydrylcarbamate derivatives were synthesized and evaluated for binding affinity to M1, M2 and M3 receptors, and for antimuscarinic activities. Receptor binding assays indicated that 1-benzyl-4-piperidyl benzhydrylcarbamate derivatives showed higher affinities for M1 and M3 receptors, and good selectivities for M3 over M2 receptor, than the corresponding ester analog. These results indicate that the urethane bond is a novel linker for muscarinic antagonists, and serves to lock the molecular conformation and allows the hydrophobic portion and cationic site of the molecule to bind to M1 and M3 muscarinic receptors. Among the prepared compounds, 1-(4-methylaminobenzyl)-4-piperidyl benzhydrylcarbamate monohydrochloride (18b, NY-58790) exhibited potent inhibitory activity on bladdar pressure in reflexly-evoked rhythmic contraction, comparable to oxybutynin and was approximately ten times less inhibitory on oxotremorine induced salivary secretion than oxybutynin in rats. Further evaluation of antimuscarinic effects on bradycardia and pressor in pithed rats, and on tremor in mice, demonstrated that VM-58790 can be useful for treatment of urinary urge incontinence as a bladder-selective M3 antagonist with

L12 ANSWER 5 OF 121 ACCESSION NUMBER:

DOCUMENT NUMBER:

CORPORATE SOURCE:

1 MEDLINE on STN
1998334184 MEDLINE
PubMed ID: 9671109
Comparison of the in vitro and in vivo profiles of
tolterodine with those of subtype-selective
mmacarinic receptor antagonists.
Gillberg P G; Sundquist S; Nilvebrant L
Department of Pharmacology, Pharmacia and Upjohn, Uppsala,
Sweden. Oweuen. European journal of pharmacology, (1998 May 22) 349 (2-3) 285-92. SOURCE:

Journal code: 1254354. ISSN: 0014-2999.

PUB. COUNTRY:

DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

285-92.
JOURNAY: Netherlands
UNENT TYPE: Journal, Article; (JOURNAL ARTICLE)
GUAGE: English
E SEGMENT: Priority Journals
RY MONTH: 199809
RY DATE: Entered STM: 19981006
Entered Medline: 1998092
TOlterodine ([R]-N,N-diisopropyl-3-(2-hydroxy-5 methylphenyl)-3-phenylpropanamine | is a new potent and competitive mascarinic
receptor antagonist developed for the treatment of urinary urge incontinence and other symptoms of overactive bladder. In vivo, tolterodine exhibits functional selectivity for the urinary bladder over salivary glands, a profile that cannot be explained in terms of selectivity for a single mascarinic receptor subtype. The alm of this study was to compare the in vitro and in vivo antimuscarinic profiles of tolterodine with those of mascarinic receptor antagonists with distinct receptor subtype-selectivity profiles: darlemacin ([S]-2-[1-[2-(2,3] dihydrobenzofuran-5-[9-1]ethyl] 3 pyrrolidinyl]-2-d iphenylacetamide; selective for muscarinic mascarinic profiles: darlemacin ([S]-2-[1-[2-(2,3] dihydrobenzofuran-5-[9-1]ethyl] 3 pyrrolidinyl]-2-d iphenylacetamide; selective for muscarinic mascarinic mascarinic

SOURCE:

ANSWER 7 OF 121 MEDLINE on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

AUTHOR:

CORPORATE SOURCE:

1 MEDLINE on STN
1998288631 MEDLINE
PubMed ID: 9626942
Prejunctional facilitatory and inhibitory
modulation of parasympathetic nerve transmission in the
rabbit urinary bladder.
Tobin G; Sjogren C
Department of Pharmacology, Institute of Physiology and
Pharmacology, University of Goteborg, Sweden..
qunnar.tobinsodontologi.qu.se
Journal of the autonomic nervous system, (1998 Feb 5) 68
(3) 153-6.

Journal code: 8003419. ISSN: 0165-1838. Netherlands

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: Journal; Article; (JOURNAL ARTICLE)

English Priority Journals

ESEGMENT: Priority Journals

REY MONTH: 199808

REY DATE: Entered STN: 19980820

Entered Medline: 19980813

Release of [3H] choline and muscle contraction in response to electrical field stimulation were measured from rabbit detrusor muscle strips previously loaded with [3H] choline. The importance of different stimulation frequencies (1 and 10 Hz) for activating either facilitatory or inhibitory prejunctional effects was examined in the presence of muscarinic and adrenergic (alpha2) receptor selective substances. At 1 Hz, neither [3H] choline overflow nor contraction was affected by the Mi-selective receptor antagonist pirenzepine (10(-7) M), whereas overflow and contraction decreased at 10 Hz. The Mi-selective receptor agonist McN-A-343 (10(-6) M) caused no significant changes except for reducing contractions at 10 Hz. The M2-selective receptor antagonist methoctramine (10(-6) M), on the other hand, increased overflow as well as contraction at both frequencies, most conspicuously at 1 Hz. Airopine (10(-7) M) caused a significant increase with respect to overflow only at 1 Hz, while quite the opposite effect occurred with respect to contractions (reduced only at 10 Hz). Clonidine (10(-6) M) induced inhibition of [3H] choline overflow at 10 Hz only, but without significantly changing contractile responses. The results show that in the rabbit urinary bladder a muscarinic autoreceptor mediated inhibition (M2) of the transmitter release dominates during low frequency stimulation and that a facilitation (M1) may be present at stimulations with higher frequencies. However, this amplification may also be influenced by alpha2 adrenoceptor mediated inhibition.

L12 ANSWER 6 OF 121
ACCESSION NUMBER: 1
DOCUMENT NUMBER: P
TITLE: 1 MEDLINE on STN 1998332162 MEDLINE PubMed ID: 9669499

AUTHOR

CORPORATE SOURCE:

PubMed ID: 9689499 Evidence for purinergic neurotransmission in the urinary bladder of pithed rats. Hegde S; Mandel D A; Wilford M R; Briaud S; Ford A P; Eglen R M Center for Biological Research, Roche Bioscience, Palo Alto, CA 94304, USA. sharath.hegde@roche.com European journal of pharmacology. (1998 May 15) 349 (1) 75-82. SOURCE:

75-62. Journal code: 1254354, ISSN: 0014-2999. Netherlands

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE: English FILE SEGMENT:

Priority Journals ENTRY MONTH: ENTRY DATE:

L12 ANSWER 8 OF 121 ACCESSION NUMBER: MEDLINE on STN

1998287656 MED PubMed ID: 9624560 MEDLINE DOCUMENT NUMBER:

TITLE:

AUTHOR:

CORPORATE SOURCE:

SOURCE:

PubMed ID: 9624560
Pharmacologic actions of temiverine (p-INN) and its active
metabolite, RCC 36, on isolated human urinary
bladder muscle.

Kikukawa H; Yoshida M; Wada Y; Nishi K; Ueda S
Department of Urology, Kumamoto University School of
Medicine, Japan.
International journal of urology: official journal of the
Japanese Urological Association, (1998 May) 5 (3) 268-75.
Journal code: 9440237. ISSN: 0919-8172.

Japan
Journal; Article; (JOURNAL ARTICLE) PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

Journal code: 9440237. ISSN: 0919-8172.

JOURNTY:
JOURNAY:
JOURNAY

L12 ANSWER 9 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

CORPORATE SOURCE:

1 MEDLINE on STN
1998280590 MEDLINE
PubMed ID: 9617596
Tolterodine.
Hills C J; Winter S A; Balfour J A
Adis International Limited, Chester, England and Auckland,
New Zealand.

PUB. COUNTRY

DOCUMENT TYPE

LANGUAGE:

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

Addis International Limited, Chester, England and Auckland, New Zealand.

New Zealand.

CC:

Drugs, (1998 Jun) 55 (6) 813-20; discussion 821-2. Ref: 39 Journal code: 7600076. ISSN: 0012-6667.

COUNTRY:

New Zealand

MENT TYPE:

JOURNAL ARTICLE)

General Review, (REVIEW)

(REVIEW, TUTORIAL)

JUAGE:

English

SEGMENT:

Priority Journals

19809

RY DATE:

Entered STN: 19980925

Last Updated on STN: 19980925

Last Updated on STN: 19980914

Tolterodine is a competitive muscarinic receptor antagonist which has recently been launched for the treatment of overactive bladder. Tolterodine shows functional selectivity for the bladder over the salivary glands in vivo, which is not attributable to muscarinic receptor subtype selectivity. It is as potent as oxybutynin in inhibiting bladder contraction, but is much less potent in inhibiting salivation, suggesting that it may have less propensity to cause dry mouth in clinical use. In patients with overactive bladder, toleterodine singinificantly reduces the frequency of micturition and number of incontinence episodes, while increasing the average volume voided. The onset of pharmacological action of tolterodine is *1 hour and therapeutic efficacy is maintained during long term treatment. In comparative trials, tolterodine is significantly better tolerated than oxybutynin over the salivation of tolterodine is *1 hour and therapeutic efficacy is maintained during long term treatment. In comparative trials, tolterodine is significantly better tolerated than oxybutynin yearticularly with respect to the incidence and severity of dry mouth. No clinically relevant ECG changes have been noted with tolterodine.

L12 ANSWER 11 OF 121 MEDLINE ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
PuBMed ID: 9485312
Prejunctional M1 facilitory and M2 inhibitory
muscarinic receptors mediate rat bladder
contractility.
AUTHOR:
CORPORATE SOURCE:
CONTRACT NUMBER:
R01-DK43333
SOURCE:

MEDLINE ON STN
1 facilitory and M2 inhibitory
muscarinic receptors mediate rat bladder
contractility.
Fraverman A S; Kohn I J; Luthin G R; Ruggieri M R
Department of Urology, Temple University School of
Medicine, Philadelphia 19140, USA.
ROI-DK43333
SOURCE:
American igures

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH

ENTRY DATE:

Medicine, Philadelphia 19140, USA.

RO1-DK493333 (NIDDK)

RO1-DK43333 (NIDK)

L12 ANSWER 10 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

AMDLINE On STN
1998178614 MEDLINE
PubMed ID: 9519799
Characterization of muscarinic receptors
mediating the contraction of the urinary detrusor
muscle in cynomolgus monkeys and guinea pigs.
Laif FM, Cobuzzi A; Spinelli W
Cardiovascular/Metabolic Diseases, Wyeth-Ayerst Research,
Princeton, NJ 08543-8000, USA.
Life sciences, (1998) 62 (13) 1179 86.
Journal code: 0375521. ISSN: 0024-3205.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English

AUTHOR: CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

English Priority Journals FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

ENGLER: Priority Journals

RY MONTH: 193804

RY MONTH: 193804

YD DATE: Entered STN: 19980416
Entered Medline: 19980406

We have characterized in vitro the muscarinic receptors
mediating the contraction of the detrusor muscle in Cynomolgus monkeys and
guinea pigs using carbachol as the agonist and 4-diphenylacetoxy-Nmethylpiperidine methiodide (4-DAMP, M3-selective), methoctramine

(M2 selective) and pirenzepine (M1 selective) as the antagonists.
Carbachol induced a concentration-dependent contraction of the detrusor
muscle of monkey and guinea pig yielding similar pD2 values of 6.574/-0.03

(n=50) and 6.774/-0.06 (n=36), respectively. In the detrusor muscle of
Cynomolgus monkey, all antagonists produced a concentration-dependent
Anhibition of carbachol-induced contractions, without decreasing
the maximal response. Schild plot analysis yielded slopes not different
from unity for all antagonists. The order of antagonist potency was:

4 DAMP (pA2-8.96)-pirenzepine (pA2-8.66)-methoctramine (pA2-6.03),
suggesting that M3 receptors have a dominant role in mediating detrusor
contraction. In the detrusor muscle of the guinea pig. 4-DAMP and
pirenzepine, but not methoctramine, produced a concentration dependent
lambition of the carbachol induced contractions, without
decreasing the maximal response. Schild plot analysis yielded a slope not
different from unity for 4-DAMP and pirenzepine. 4-DAMP (pA2-9.07) had a
higher potency than pirenzepine (pA2-6.66), a finding consistent with
previously published data. The present study shows that in Cynomolgus
monkey stimulation of the M3 subtype is dominant in mediating detrusor
contraction upon carbachol stimulation.

L12 ANSWER 12 OF 121 ACCESSION NUMBER: 1 DOCUMENT NUMBER:

21 MEDLINE on STN
1998136057 MEDLINE
PubMed ID: 9477190
Nuscarinic receptor subtypes and receptor coupled
phosphatidylinositol hydrolysis in rat bladder
smooth muscle. TITLE:

Mimata H; Nomura Y; Emoto A; Latifpour J; Wheeler M; Weiss AUTHOR:

R M Department of Urology, Oita Medical University, Japan. International journal of urology: official journal of the Japanese Urological Association, (1997 Nov) 4 (6) 591-6. Journal code: 9440237. ISSN: 0919-8172. CORPORATE SOURCE: SOURCE:

PUB. COUNTRY:

Japan Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE:

English Priority Journals 199804 FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEMMENT: Priority Journals

Y MONTH: 199804

Entered STN: 19980416

Entered Medline: 19980408

BACKGROUND: The purpose of this study was to evaluate the muscarinic receptor subtypes expressed in rat bladder smooth muscle and characterize the muscarinic receptor-coupled phosphatidylinositol (PI) hydrolysis in order to clarify the first step of bladder smooth muscle contraction. METHODS: Expressions of mRNAs of muscarinic receptor subtypes were examined by Northern blot analysis. Changes in the mass of inositol 1.4.5-triephosphate (IP3) and the inhibitory effects of muscarinic subtype specific antagonists on PI hydrolysis were determined after carbachol stimulation. RESULTS: mRNAs of m2 and m3 genes, encoding M2 and M3 receptors, were expressed in rat bladder smooth muscle. Carbachol produced a rapid increase of IP1, which returned to the basal level within 30 seconds. 4-Diphenylacetoxyl N-methylpiperidine methiodide (4-DAMP; M1 and M3 antagonist) strongly inhibited the PI hydrolysis, but methoctramine (M2 antagonist) partially inhibited it at 10(-4) mol/L. The ICSO value for acropine was \$5.5 x 10(-9) mol/L, for pirenzepine 6.4 x 10(6) mol/L, and for 4-DAMP 1.5 x 10(-7) mol/L. CONCLUSION M2 and M3 receptors are expressed in rat wrinary bladder. Only M3 receptors are expressed in rat wrinary bladder. Only M3 receptor was involved in the production of IP3, which might induce the initial phase of contractile response in rat bladder smooth muscle after carbachol stimulation.

L12 ANSWER 13 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

CORPORATE SOURCE:

21 MEDLINE on STN
1998118820 MEDLINE
PUBMed ID: 9457494
Effect of choline ester analogues, noradrenaline and
nifedipline on normal and hypertrophied human
urinary bladder detrusor muscle.
King J A; Huddart H; Staff W G
Division of Biological Sciences, Lancuster University, UK.
General pharmacology, (1998 Jan) 30 (1) 131-6.
Journal code: 7602417. ISSN: 0306-3623.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English SOURCE

PUB. COUNTRY

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH

English
Priority Journals ENTRY DATE:

SIMORE: English
Priority Journals
SEMOMENT: Priority Journals
Finered STN: 19980319
Last Updated on STN: 19980319
1. Acetylcholine, bethanechol, carbachol and propionylcholine were all agonists of normal human detrusor smooth muscle. The order of potency was found to be carbachol > acetylcholine> bethanachol > propionylcholine. 2. In hypertrophied detrusor preparations were less sensitive to carbachol ine, but hypertrophied detrusor preparations were less sensitive to carbachol than normal detrusor smooth muscle. 3. Noradremaline had no direct effect on either normal or hypertrophied detrusor muscle, but it had a reversible inhibitory effect on the spontaneous contractile activity of normal detrusor preparations. Hypertrophied detrusor preparations waslly lacked such spontaneous activity. 4. In calcium-free saline, agonist-induced responses of both normal and hypertrophied detrusor muscle were dramatically reduced indicating that choline ester activity in the muscles was strongly dependent upon extracellular calcium. 5. Nifedipine at 10(-5) mol 1-1 inhibited acetylcholine responses and K(+)-induced contractures of both normal and hypertrophied detrusor muscles. Acetylcholine-induced responses of normal detrusor preparations were much more sensitive to inhibition by infedipine than were the responses of hypertrophied detrusor muscle. 6. The properties and densities of both the muscarinic cholinoreceptors and calcium channels appear to have been altered by the hypertrophic response secondary to benign prostatic hyperplasia.

ANSWER 15 OF 121 L12 ANSWER 15 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

21 MEDLINE on STM 1998075781 MEDLINE PubMed ID: 9413860 Subcellular distribution of SERCA and calcium-activated

AUTHOR:

Subcellular distribution of SERCA and calcium-activated ATPase in rabbit and human urinary bladder smooth muscle. Levin R M; Nicholas T J; Snithoff G G; Mandell J; Russell D; Wilbur H J; Mogavero L J Department of Biological Science, Albany College of Pharmacy, NY 12208 USA. IROL DK 26508 (NIDD). Pharmacology, (1997 Dec) 55 (6) 309-16. Journal code: 0152016. ISSN: 0031-7012. Switzerland CORPORATE SOURCE:

CONTRACT NUMBER: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: Journal; Article; (JOURNAL ARTICLE) English

FILE SEGMENT: ENTRY MONTH:

COMENT: TYPE: Journal; Article; (JOURNAL ARTICLE)

GUAGE: English

E SEGMENT: Priority Journals

RY MONTH: 19980

RY DATE: Entered Stn: 19980217

Last Updated on STN: 19980217

Entered Medline: 199802017

Frevious studies have demonstrated that calcium storage and release from IP3-dependent sites in the sarcoplasmic reticulum play an important role in the contractile response of the rabbit urinary

bladder to both field stimulation (mediated via neurotransmitter release) and bethanchol (direct muscarinic stimulation). In view of the importance of SERCA (see text) in urinary

bladder smooth muscle function, we studied the distribution of SERCA by two methods; using Western blotting to quantitate the protein concentration and by enzyme analysis using thapsigargin to specifically inhibit SERCA. Rabbit and human samples of urinary

bladder smooth muscle were homogenized and the homogenate separate into three particulate fractions by different centrifugation; the cell wall-nuclear, mitochondrial, and microsomal. The protein concentration of these three particulate fractions was determined and the SERCA protein level quantitated by Western blotting using SERCA-2 antibodies. The calcium ATPase activity was quantitated using standard enzymatic analysis and the thapsigargin sensitivity determined. The results demonstrated that (1) the concentration of SERCA was significantly greater in the microsomal fraction than in either of the other fractions for both rabbit and human bladder smooth muscle; (2) the enzymatic activities of both total calcium activated ATPase and that carcivity of both total calcium activated ATPase and the properties of the rabbit exceeded that of the human. In conclusion, the distribution of SERCA and calcium ATPase of the rabbit exceeded that of the human. In conclusion, the distribution of SERCA and calcium ATPase of the rabbit exceeded that of the human. In conclusion, the distribution of SERCA and calcium ATPase of the rabbit exceeded that of the human. In conclusion, the distribution of SERCA a

L12 ANSWER 14 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 1998088162 MEDLINE
DOCUMENT NUMBER: 191808162 MEDLINE
TITLE: The overactive bladder: pharmacologic basis of

drug treatment.

drug treatment.
Andersson K E
Department of Clinical Pharmacology, Lund University
Hospital, Sweden.
Urology, (1997 Dec) 50 (6A Suppl) 74-84; discussion 85-9.
Ref: 107
Journal code: 0366151. ISSN: 0090 4295.
United Statee
Journal; Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
(REVLEW, ACADEMIC) AUTHOR: CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

English

LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: Digitsh Priority Journals 199801

English
E SEGMENT: Priority Journals
EY MONTH: 199801
EXT MONTH: 199801
EAST Updated on STN: 19980206
Last Updated on STN: 19980206
Entered Meddine: 19980129
CBJECTIVES: To provide an overview of the basis for drug treatment of the overactive bladder. METHODS: Dublished information is evaluated. RESULTS: The causes of bladder overactivity are not known, but theoretically, increased afferent activity, decreased inhibitory control in the central nervous system (CNS) or peripheral ganglia, and increased sensitivity of the detrusor to efferent stimulation may be involved. Several CNS transmitters can modulate voiding, but few useful drugs with a defined CNS site of action have been developed. Drugs that stimulate gamma-aminobutyric acid receptors are used clinically. Potentially, drugs affecting opioid. 5-hydroxytryptamine, norepinephrine, dopamine, and glutamatergic receptors and mechanisms can be developed, but a selective action on the lower urinary tract may be difficult to obtain. Traditionally, drugs used for treatment of bladder overactivity have had a peripheral site of action, mainly efferent neurotransmission or the detrusor itself. Antimuscarinic drugs, beta-adrenoceptor agonists, drugs affecting membrane channels, patha-adrenoceptor antagonists, drugs affecting membrane channels, postaglandin synthetase inhibitors, and several other agents have been used with limited success. New information on the alpha-adrenoceptor and mascarinic receptors in the human detrusor has emerged and may be the basis for the development of new compounds with effects on bladder overactivity. Decreasing afferent activity seems an attractive therapeutic approach, and drugs affecting afferent nerves by causing release of tachykinins, such as capsaicin and analogs, as well as agents blocking tachykinin receptors, may be of therapeutic interest.

L12 ANSWER 16 OF 121
ACCESSION NUMBER: 1
DOCUMENT NUMBER: POTITLE: BO

21 MEDLINE on STN
1998063137 MEDLINE
PubMed ID: 9400490
Bethanechol activates a post receptor negative feedback
mechanism in rabbit urinary bladder

AUTHOR: CORPORATE SOURCE:

mechanism in rabbit urinary bladder smooth muscle. Shenfeld O Z; Morgan C W; Ratz P H Department of Urology, Eastern Virginia Medical School, Norfolk 23501, USA. Journal of urology, (1998 Jan) 159 (1) 252-7. Journal code: 0376374. ISSN: 0022 5347. United States SOURCE:

PUB. COUNTRY

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: Journal; Article; (JOURNAL ARTICLE) English

Abridged Index Medicus Journals; Priority Journals 199712 ENTRY MONTH: ENTRY DATE:

GUAGE:
ESEMENT: Abridged Index Medicus Journals; Priority Journals
RY MONTH: 199712
RY DATE: Entered STN: 19980116
Entered Medicus 1997122
PURPOSE: Recent studies using vascular and gut smooth muscles indicate
that contractile receptor agonists may activate post-receptor
down-regulatory mechanisms causing a temporary reduction in the strength
of subsequent contractions. Our data indicate a similar mechanism exists
in detrusor smooth muscle of the urinary bladder.
MATERIALS AND METHODS: Each isolated strip of female rabbit detrusor was
placed in a tissue bath, secured to an isometric force transducer, and
length-adjusted until depolarization with 110 mM KCl produced a maximum
contraction (S0). Subsequent contractions were normalized to S0 (S/S0) or
to a first stimulus with 30 mM KCl or caffeine (S/S1). Tissues were
pretreated with the muscarinic receptor agonist, bethanechol
(SB), then stimulated with KCl, caffeine, or Bay k 8644 to identify
potential post-receptor down regulation. RESULTS: Contractions induced by
30 mM KCl had three phases labeled fast peak (FP), slow peak (SP) and
stready state (SS). In tissues exposed for 30 min. to a maximum BE
concentration then washed for 5 min., the KCl-induced FP and SF, but not
SS, responses were reduced by approximately 40%. Smaller reductions in
peak KCl-induced contractions occurred in tissues pretreated for a shorter
duration or with a 100-fold lower BE concentration. This down-regulation
induced by bethanechol pretreatment was reversible, lasting approximately
1-2 h. Not only were KCl-induced contractions reduced by BE pretreatment,
but also those produced by the intracellular Ca(2+)-mobilizer, caffeine,
and the L+type Ca2+ channel agonist, Bay 8644. CONCLUSIONS:
Pretreatment of isolated strips of rabbit detrusor with a
muscarinia receptor agonist produced short-term down-regulation of
both influx of extracellular Ca2+ and release of intracellular Ca2+.
Reductions in the degree of this novel modulatory response during disease
conditions and aging could enhance cont

L12 ANSWER 17 OF 121 ACCESSION NUMBER: 1 MEDLINE on STN 1998049688 MEDLINE PubMed ID: 9388365 DOCUMENT NUMBER:

CORPORATE SOURCE:

runmed ID: 9388365 Autoradiographic localization of muscarinic receptors in diabetic rat bladder. Saito M; Nakamura I; Miyagawa I Department of Urology, Tottori University School of Medicine.

SOURCE:

Medicine. Nippon Hinyokika Gakkai zasshi, japanese journal of urology, (1997 Oct) 88 (10) 858-67. Journal code: 2984841R. ISSN: 0021-5287.

PUB. COUNTRY:

Journal code. DJJ Japan Journal; Article; (JOURNAL ARTICLE) Japanese Priority Journals DOCUMENT TYPE:

LANGUAGE:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals

87 MONTH: 199712

87 MONTH: 199712

87 MONTH: Entered STN: 19980122

Entered Medline: 19971231

FURPOSE: We evaluated the alterations of density, localization and subtype specificity of muscarinic receptors in experimentally-induced diabetic rat bladder. METHODS: Five groups of rats were maintained for sixteen weeks 1) diabetic, 2) diabetic insulin-treated (insulin started 8 weeks after the onset of diabetes), 3) sucrose-fed diuretic, 4) sucrose-removed and 5) age matched controls. We used radioligand binding technique and light microscopic autoradiography to define the density and distribution of muscarinic receptors in the rat urinary bladder. RESULTS & CONCLUSION:

Saturation experiments showed an increase in the density of muscarinic receptors in the bladders of diabetic and diuretic rats compared with age matched controls. Insulin treatment partially reversed the up-regulatin of muscarinic receptors in rat urinary bladder. Autoradiographic studies also indicated that muscarinic receptors were located in all layers of the bladder muscularis. The muscularis of the bladder dome contained higher densities of muscarinic receptors than that of the bladder base, Lack of [3H]ONB binding to transitional epithelium, lamina propria, and tunica adventitia suggests an absence of muscarinic receptors in these regions. Inhibition of [3H]ONB binding to the bladder sections by selective muscarinic antagonists indicated existence of the M2 and M3 receptor subtypes in the muscularis of the rat bladder.

ANSWER 19 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR CORPORATE SOURCE:

21 MEDLINE On STN
1998015289 MEDLINE
PubMed ID: 9353847
Antimuscarinic potency and bladder selectivity of
PNU-200577, a major metabolite of tolterodine.
Nilvebrant L; Gillberg P G; Sparf B
Medical Department Urology, Pharmacia & Upjohn AB, Uppsala,
Sweden.

Sweden.
Pharmacology & toxicology, (1997 Oct) 81 (4) 169-72.
Journal code: 8702180. ISSN: 0901-9928. SOURCE:

Denmark Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

English FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
The priority Jour

L12 ANSWER 18 OF 121
ACCESSION NUMBER: 19
DOCUMENT NUMBER: PU
TITLE:

21 MEDLINE on STN
1998019306 MEDLINE
PubMed ID: 9353395
MR muscarinta autoreceptor mediated
inhibition of -3H-acetylcholine release in the rat
isolated urinary bladder.
D'Agostino G; Barbieri A; Chiossa E; Tonini M
Institute of Pharmacology, School of Pharmacy, University
of Payia, Pavia, Italy.
Journal of pharmacology and experimental therapeutics,
(1997 Nov) 283 (2) 750 6.
Journal code: 0376362. ISSN: 0022-3565.
United States
Journal, Article; (JOURNAL ARTICLE)
English AUTHOR: CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: English Priority Journals 199712

GUAGE: English E SEDMENT: Priority Journals RX MONTH: 199712

RX MONTH: 199712

RX DATE: Entered SIN: 19980109

Last Updated on SIN: 19980109

Last Updated on SIN: 19980109

A pharmacological analysis was carried out in the rat urinary bladder to assess the nature of muscarinic receptors subtypes functionally involved in the negative feedback mechanism regulating acetylcholine (ACh) secretion from postganglionic cholinergic nerve terminals and in smooth muscle contraction. Bladder strips were preincubated with 3H choline, and the electrically evoked (JH)ACh release was detected simultaneously with contraction in the absence of acetylcholinesterase inhibitors. The effects were compared of seven muscarinic antagonists on [3H]ACh secretion (prejunctional effect) and muscle contraction (postjunctional effect). The rank order of postjunctional potencies (-log EGO) for the seven antagonists (atropine > 4-diphenylacetoxy-N-methylpiperidine methiodide (4-DAMP) hexahydrosiladiphenidol hydrochloride (HRSID) > tripitramine > pirenzepine > AF DX.-116 > methoctramine) as well as their postjunctional affinity estimates (PA2) are in keeping with the notion that muscarinic receptors responsible for bladder contraction belong to the M3 subtype. The M3 subtype-perferring 4-DAMP and HHSID did not discriminate between prejunctional and postjunctional effects. The M3/M4 subtype-preferring antagonists tripitramine, methoctramine and AF-DX 116 were more potent in facilitating the evoked [3H]ACh release than in inhibiting the contractile response. The rank order of prejunctional potencies was atropine > 4-DAMP > tripitramine > HHSID > methoctramine > AF-DX 116 > pirenzepine, indicating the involvement of M4 receptors. Furthermore, when potency relationship was determined by correlations were significant for both M4 and M5 subtypes, but the best correlation found (P < 001) was for the M4 subtype. These findings suggest that the negative feedback mechanism inhibiting the

21 MEDLINE on STN
1998006873 MEDLINE
PUMMed TD: 9247324
Anticholinergic and calcium antagonistic activities of
NS-21 contribute to the inhibition of rat
urinary bladder contractions
Hamada K; Sasaki Y; Taniguchi N; Fukui H; Miyatsuka Y;
Kimura Y; Ukai Y; Yoshikuni Y; Kimura K
Research laboratories, Nippon Shinyaku Co., Ltd., Kyoto,
Jaoan. L12 ANSWER 20 OF 121
ACCESSION NUMBER: 10
DOCUMENT NUMBER: PATITLE: A

CORPORATE SOURCE:

Japan.
General pharmacology, (1997 Nov) 29 (5) 771-8.
Journal code: 7602417. ISSN: 0306-3623.
EWGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

AUGUSTATION CONTRACT OF THE MAIN ARTICLE)

LOUIS RESIDENCE:

LOUIS SERVINE:

L

L12 ANSWER 21 OF 121 ACCESSION NUMBER: 1 MEDLINE on STN DOCUMENT NUMBER:

21 MEDLINE on STN
1998004388 MEDLINE
PUBMed ID: 9346402
Subcellular distribution of SERCA and calcium-activated
ATFase in rabbit and human urinary
bladder smooth muscle.
Levin R M, Nicholas T J; Snitkoff G G; Mandell J; Russell
D; Wilbur H J; Mogavero L J
Albany College of Pharmacy, Stratton Veterans Affairs
Medical Center, N.Y. 12208, USA.
1ROI DX 26508 (NIDDK)
Pharmacology, (1997 Sep) 55 (3) 136-43.
Journal code: 0152016. ISSN: 0031-7012.
Switzerland AUTHOR

CORPORATE SOURCE:

CONTRACT NUMBER: SOURCE:

PUB. COUNTRY:

Switzerland
Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: English

FILE SEGMENT: Priority Journals 199711

ENTRY MONTH:

ENTRY DATE:

ESCRENT: Priority Journals
RY MONTH: 199712
Last Updated on STN: 19971224
Entered Medline: 19971119
Previous Studies have demonstrated that calcium storage and release from 19-3-dependent sites in the sarcoplasmic reticulum play an important role in the contractile response of the rabbit urinary
bladder to both field stimulation (mediated via neurotransmitter release) and bethanechol (direct muscarinic stimulation). In view of the importance of SERCA in urinary bladder smooth muscle function, we studied the distribution of SERCA by two methods: using Western blotting to quantitate the protein concentration and by enzyme analysis using thapsigargin to specifically inhibit SERCA. Rabbit and human samples of urinary bladder smooth muscle were homogenized and the homogenize separated into three particulate fractions by differential centrifugation: nuclear-cell wall, mitochondrial, and microsomal. The protein concentration of these three particulate fractions was determined and the SERCA protein level quantitated by Western blotting using SERCA-2 antibodies. The calcium-ATPase activity was quantitated using standard enzymatic analysis and the thapsigargin sensitivity determined. The results demonstrated that: (1) the concentration of SERCA was significantly greater in the microsomal fraction than in either of the other fractions for both rabbit and human bladder smooth muscle; (2) the enzymatic activities of both total calcium-activated ATPase and thapsigargin-sensitive calcium ATPase were evenly divided among the three fractions, and (3) the enzymatic activity of both total calcium-activated ATPase and thapsigargin-sensitive calcium ATPase of the rabbit bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to that in the human bladder smooth muscle was similar to t

ANSWER 23 OF 121 21 MEDLINE on STN 97308396 MEDLINE ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

MEDLINE On STN
97308396 MEDLINE
PubMed ID: 9165622
Etiology of bladder dysfunction secondary to
partial outlet obstruction. Calcium disregulation in
bladder power generation and the ability to perform

AUTHOR:

work. Levin R M; Yu H J; Kim K B; Longhurst P A; Wein A J; Damaser M S Division of Urology, University of Pennsylvania, USA. 5 T32-DK07708 (NIDDK)

CORPORATE SOURCE:

CONTRACT NUMBER: 5 T32 RO1 DK 46508 (NIDDK) SOURCE:

NIDDK)
Scandinavian journal of urology and nephrology.
Supplementum, (1997) 184 43-55.
Journal code: 0153034. ISSN: 0300 8886.

Sweden

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: Sweden Journal; Article; (JOURNAL ARTICLE) English Priority Journals 199708

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

L12 ANSWER 22 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 97344033 MEDLINE
DOCUMENT NUMBER: 9200560
TITLE: Pubmed ID: 9200560
Tolterodine- a new bladder-selective
antimuscarinic agent.
AUTHOR: AUTHOR: SOURCE: Selective antimuscarinic agent.
Milvebrant L; Andersson K E; Gillberg P G; Stahl M; Sparf E
Medical Department Urology, Pharmacia & Upjohn AB, Uppsala,
Sweden.

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

antimuscarinic agent.

Nalivebrant L; Andersson K E; Gillberg P G; Stahl M; Sparf E PORATE SOURCE:

Milvebrant L; Andersson K E; Gillberg P G; Stahl M; Sparf E PORATE SOURCE:

Milvebrant L; Andersson K E; Gillberg P G; Stahl M; Sparf E BORATE SOURCE:

European journal of pharmacology, (1997 May 30) 327 (2-3) 195-207.

Journal code: 1254354 ISSN: 0014 2999.

Netherlands

JUAGE:

English

E SEGMENT:

Priority Journals

RY MONTH:

199708

RY DATE:

Entered Sellish:

Tolterodine is a new mucarinic receptor antagonist intended for the treatment of urinary urge incontinence and other symptoms related to an overactive bladder. The aim of the present study was to compare the antimuscarinic properties of tolterodine with those of oxybutymin, in vitro and in vivo. Tolterodine effectively inhibited carbachol-induced contractions of isolated strips of urinary bladder from guinea pigs (K(B) 3.0 nm; pA2 8.4; Schild slope 1.04) in a concentration-dependent, competitive manner. The affinity of tolterodine was similar to that derived for oxybutymin (K(B) 4.4 nm; pA2 8.5; Schild slope 0.89) in the guinea-pig bladder. Tolterodine (21-2103 nmol/kg (0.01-1 mg/kg); intravenous infusion) was significantly more potent in inhibiting acetyleholine-induced urinary bladder contraction than electrically-induced salivation in the anaesthetised cat. In contrast, oxybutynin displayed the opposite tissue selectivity. Radioligand binding data showed that tolterodine bound with high affinity to muscarinic receptors in urinary bladder from humans (K(i) 3.3 nm). Tolterodine and oxybutynin bound with 8-times higher affinity (K(i) 0.62 nm). Binding data on human suscarinic microexperse in the paroting land, where oxybutynin bound with 8-times higher affinity (K(i) 0.62 nm). Binding data on human suscarinic microexperse in the paroting land, where oxybutynin bound with 8-times higher affinity (K(i) 2.8 and 6.7 nm) that at muscarinic M3/m3 receptors (K(i) 2.8 and 6.7 nm). The tissue selectivity demonstrated for tolterodine in vivo cannot be attributed

L12 ANSWER 24 OF 121
ACCESSION NUMBER: 9
DOCUMENT NUMBER: P
TITLE: R

MEDLINE on STN
97276408 MEDLINE
PUDMed ID: 9130161
Role of L- and N-type Ca2+ channels in muscarinic
receptor-mediated facilitation of Ach and noradrenaline
release in the rat urinary bladder.
Somogyi G T; Zernova G V; Tanowitz M; de Groat W C
Department of Pharmacology, University of Pittsburgh, PA
15261, USA. somon-%pitt.edu
NIDDR-45741 (NIDDW)
Journal of ohywiology, (1997 Mar 15) 499 (Pt 3) 645 54 AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER: SOURCE:

Journal of physiology, (1997 Mar 15) 499 (Pt 3) 645 54. Journal code: 0266262. ISSN: 0022-3751. BOGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: English

LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

MENT TYPE: Journal; Article; (JOURNAL ARTICLE)
NUAGE: English

2 SEGMENT: Priority Journals

2 Y MONTH: 199708

2 MATE: Entered STN: 19970825

Last Updated on STN: 20000303

Entered Medline: 19970812

1. 3H-Noradremaline (NA) and 14C-acetylcholine (ACh) released by
electrical field stimulation were measured simultaneously in strips from
the body of rat urinary bladder. 2 onega-Contoxin

GVIA (Omega-CgTK; 20-100 nM) suppressed the non-facilitated transmitter
release evoked by intermittent stimulation (15), whereas nifedipine (1
microM) did not affect release. 3. Continuous electrical stimulation (CS)
facilitated NA and ACh release. 3. Continuous electrical stimulation (CS)
facilitated NA and ACh release. 3. Continuous alectrical stimulation (CS)
facilitated NA and ACh release. Nifedipine depressed ACh release (434) but not NA
release. Combined administration of infedipine and omega-CgTX (20 nM)
produced a greater suppression of NA and ACh release (85 and 914,
respectively). 4. Maximal muscarinic facilitation of NA
(5-fold) and ACh (17-fold) release occurred following administration of
seerine, an anticholinesterase agent. Release of both NA and ACh was
depressed by nifedipine (70 and 83%, respectively) but not by omega-CgTX.
Combined application of omega-CgTX and nifedipine elicited a further
depression of NA (5%) but not ACh release. 5. When NA and ACh release
was facilitated with phorbol idubutysate (0.5 microM), nifedipine
inhibited ACh (67%) but not NA release, whereas omega-CgTX
inhibited NA (73%) but not ACh release. Combined administration
of both Ca2+ channel blockers did not elicit greater inhibition.
6. Bay K 8644, the L-type Ca2+ channel activator, increased ACh release
in a dose-dependent manner (up to 5-fold) but did not significantly change
NA release. 7. Both omega-CgTX (20-100 nM) and nifedipine (100 nM-1
microM) significantly decreased (50-80%) the neurally evoked contractions
of the bladder strips. 8. It is concluded that L-type Ca2+
channels play a major role in muscarinic facilitation of NA a

L12 ANSWER 25 OF 121
ACCESSION NUMBER: 91
DOCUMENT NUMBER: PETITLE: EF MEDLINE on STN

21 MEDLINE on STN
97273377 MEDLINE
97273377 MEDLINE
PUMMed ID: 9127817
Effects of ONO-2235, an aldose reductase inhibitor
on muscarinic receptors and contractile
response of the urinary bladder in rats
with streptosotocin induced diabetes.
Kanda M; Eto K; Tanabe N; Sugiyama A; Hashimoto K; Ueno A
Department of Urology, Yamanashi Medical University, Japan.
Japanese journal of pharmacology, (1997 Mar) 73 (3) 221-8.
Japan
Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals
199707
Entered STN: 19970805 AUTHOR CORPORATE SOURCE: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
IY MONTH: 199707
IY MONTH: 199707
IX MONTH: 199707
IX DATE: Entered STN: 19970805
Entered Medline: 19970721
This study was conducted to evaluate effects of the aldose reductase inhibitor ONO-2235 on the contractile response to acetylcholine of the urinary bladder dome of streptozotocin-induced diabetes mellitus (DM) rats and simultaneously observe the changes in the function and number of muscarinic receptors and the sorbitol content of the bladder. The contractile response to acetylcholine increased 51% in the DM rat bladder dome compared to the normal rats; however, this was attenuated to a 10% increase by administration of 100 mg/kg ONO-2235 for 2 weeks. Treatment with ONO-2235 significantly decreased the specific (3M] quinuclidinyl benzilate binding in DM rats. However there was no significant dose-dependency among the ONO-2235-treated groups. The sorbitol levels of the sciatic nerve and the bladder were higher in the DM rats compared to the control rats; ONO-2235 decreased the level, although it did not completely reverse them to the control level. These results suggest that an aldose reductase inhibitor attenuates the increase of the muscarinic receptor number and normalizes the enhanced contractile response to acetylcholine caused by hyperglycemia and diuresis, probably through suppression of the polyol-pathway in the DM rat bladder dome.

ANSWER 26 OF 121 MEDLINE on STN (Continued) potency (ID35%inh, nmol kg 1, i.v.): 4-DAMP (8.1) > atropine (20.7) > methoctramine (119.9) > darifenacin (283.3) > pirenzepine (369.1) > methoctramine (119.9) > darifenacin (283.3) > pirenzepine (369.1) > p F HHSIO (1053.8). These potency estimates correlated most favourably (r = 0.89, P = 0.04) with the pKi estimates of these compounds at human recombinant manacarine n2 receptors. This is consistent with a major contribution of M2 receptors in the generation of volume induced bladder contractions, although the modest potency of darifenacin does not exclude a role of M3 receptors. Fretreatment with propranolol (1 mg kg-1, i.v.) increased the ID35%inh of methoctramine significantly from 95 9 to 404.5 nmol kg-1 but had no significant effects on the inhibitory responses to darifenacin. These data suggest an obligatory role of beta adrenoceptors in M2 receptor-mediated bladder contractions in vivo. 5. The findings of the present study suggest that both M2 and M3 receptors can cause contraction of the rat bladder in vitro and may also mediate reflex bladder contractions in vivo. It is proposed that mascarinic M3 receptor activation primarily causes direct contraction of the detrusor whereas M2 receptor activation can contract the bladder indirectly by reversing sympathetically (i.e. beta-adrenoceptor)-mediated indirectly by reversing sympathetically (i.e. beta-adrenoceptor)-mediated relaxation. This dual mechanism may allow the parasympathetic nervous system, which is activated during voiding, to cause more efficient and complete emptying of the bladder.

L12 ANSWER 26 OF 121
ACCESSION NUMBER: 95
DOCUMENT NUMBER: POTITLE: MEDLINE on STN

MEDITING ON STN
97268065 MEDITINE
PubMed ID: 9113359
Functional role of M2 and M3 muscarinic receptors
in the urinary bladder of rate in vitro

and in vivo. Hegde S S; Choppin A; Bonhaus D; Briaud S; Loeb M; Moy T M; AUTHOR:

and in 11vo. Hegde S S; Choppin A; Bonhaus D; Briaud S; Loeb M; Moy Loury D; Eglen R M Department of Urogenital and Mechanistic Pharmacology, Institute of Pharmacology, Palo Alto, CA, USA. British Journal of pharmacology, (1997 Apr) 120 (8) 140c.18 CORPORATE SOURCE:

SOURCE:

1409-18.
Journal code: 7502536. ISSN: 0007-1188.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)

English

Priority Journals 199706

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

L12 ANSWER 27 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: P MEDLINE on STN

21 MEDLINE on STN
97149597 MEDLINE
PubMed ID: 8996405
Evidence for the presence of regional differences in the
subtype specificity of muscarinic receptors in
rabbit lower urinary tract.
Mutoh S; Latifpour J; Saito M; Weiss R M
Section of Urology, Yale University School of Medicine, New
Haven, Connecticut 06520-8041, USA.
DK 38311 (NIDDK) AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER

DK 42530 (NIDDK) SOURCE: J

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

Haven, Connecticut 06520-8041, USA.

TRACT NUMBER: DX 38311 (NIDDK)

RCE: Journal of urology, (1997 Feb) 157 (2) 717 21.

Journal code: 0376374. ISSN: 0022 5347.

COUNTRY: United States

UMENT TYPE: Journal, Atticle; (JOURNAL ARTICLE)

GUAGE: English

E SEDMENT: Abridged Index Medicus Journals; Priority Journals

RY MONTH: 199702

RY MONTH: 199702

RY MONTH: 1997035

Lest Updated on STN: 19970305

Lest Updated on STN: 19970305

To elucidate the subtype specificity of muscarinic cholinergic receptors in mediating contractile responses in the lower urinary tract, we investigated contractile and biochemical properties of muscarinic receptors in bladder dome, bladder

base and urethra of the rabbit. Isometric contractile response curves to increasing concentrations of carbachol were constructed in the absence and presence of various concentrations of subtype selective muscarinic antagonists. Bladder dome, bladder base, and urethra demonstrate different characteristics in terms of efficacy and potency with respect to carbachol-induced contractile responses. Emax values are significantly larger and ED50 values are significantly smaller in bladder dome and bladder base than in urethra.

Calculation of the pA2 values, the negative logarithm of the antagonist affinity constant (RB), for a series of muscarinic antagonists, i.e., atropine (nonselective), pireuzepine (MI selective), methoctramine (MZ selective), and 2 DAMF (MI/MS selective) indicate that the carbachol-induced contractile response in bladder dome and bladder hase is mediated through the MI and/or M3 and possibly M2 subtypes *Muscarinic cholinergic antagonists inhibit (BH quinulidiny) benzilate cholinergic antagonists inhibit (BH quinulidiny) benzilate cholinergine. The binding data indicate the predominance of the M2 preseptors subtype in all three regions.

L12 ANSWER 28 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 97144729 MEDLINE
DOCUMENT NUMBER: PubMed ID: 8990491
TITLE: AUTHOR: CORPORATE SOURCE: Hass M A; Geloso D; Leggett R E; Horan P; Levin R M
CONTRACT NUMBER: Dk-26508-15 (NIDDK)
CONTRACT NUMBER: DK-26508-15 (NIDDK)
FARMACOLOGY (1996 Nov) 53 (5) 320-7.

Pharmacology, (1996 Nov) 53 (5) 320-7. Journal code: 0152016. ISSN: 0031-7012. Switzerland Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: English

Priority Journals 199706 Entered STN: 19970620 Last Updated on STN: 19990129 Entered Medline: 19970612

Last Updated on STN: 19970620

Last Updated on STN: 19990129

Entered Medline: 19970612

Urinary bladder smooth muscle contraction can be evaluated using field stimulation (neurohumoral transmission), carbachol (muscarinic stimulation), and KCl (direct membrane depolarization).

We recently evaluated the activity of a novel organic chemical, macrocycle-1, on the contractile responses of the bladder to field stimulation, carbachol, and KCl. Isolated strips of rabbit bladder were mounted in individual baths containing 7.5 ml Tyrode's solution. The response to FS (1-32 Hz), carbachol (1 mumol/1), and KCl (120 mmol/1) were determined in the presence and absence of 3 different concentrations of macrocycle-1. Maximal tension, the rate of tension generation, the time to maximal tension, and the rate of decay following maximal tension on were determined. The results can be summarized as follows: (1) In the absence of macrocycle-1, maximal tension and the maximal and mean rate of tension generation increased with frequency, whereas the time to maximal tension was constant. The rate of decay of tension following maximal tension was greater for 8, 16 and 32 Ha as compared to 1 or 2 Hz. (2) The maximal response to KCl was lower than either FS or carbachol. The maximal rates of tension generation for KCl was greater than that of FS, and the rate of tension generation for KCl was greater than that of either carbachol or FS. (3) Macrocycle-1 had a greater inhibitory effect on KCl stimulation than on carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation; and a greater inhibitory effect on KCl and carbachol stimulation than on FS. (4) The rate of tension generation was more sensitive to macrocycle-1 is acting as an intracellular calciu

AUTHOR: CORPORATE SOURCE:

L12 ANSWER 30 OF 121 MEDLINE ON STN
ACCRSSION NUMBER: 97066736 MEDLINE
DOCUMENT NUMBER: Pubmed ID: 9810212
TITLE: 910mscarinic receptor-induced facilitation of

M1 muscarinic receptor-induced facilitation of ACh and noradrenaline release in the rat bladder is mediated by protein kinase C. Somogyi G T; Tanowitz M; Zernova G; de Groat W C Department of Pharmacology, University of Pittsburgh, PA 15261, USA. somoe@pitt.edu DK-45741 (NIDDK) Journal of physiology, (1996 Oct 1) 496 (Pt 1) 245-54. Journal code: 0266262. ISSN: 0022-3751. ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE)

CONTRACT NUMBER: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE; LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

.COUNTRY: ENCLAND: United Kingdom MENT TYPE: Journal; Article; (JOURNAL ARTICLE)
RUAGE: English
ESEGMENT: Priority Journals
RY MONTH: 199703
RY DATE: Entered STN: 19970327
Last Updated on STN: 19970327
Entered Medline: 19970314
1. [3H] Noradrenaline (NA) AND [14C]acetylcholine (ACh) released by electrical field stimulation were measured simultaneously in strips from the body of rat urinary bladder. 2. [3H]NA and [14C]ACh release was greater during continuous stimulation (CS; 10 Hz, 100 shocks) or in the presence of eserine than during intermittent train stimulation (IS; 10 Hz, 10 shocks every 5 s, 10 times). Atropine (1 macroM) or pirensepine (0.05-0.1 microM) blocked the CS- or eserine facilitated release. 3. The protein kinase C (PKC) activator phorbol dibutyrate (PDB; 0.05 and 0.5 microM) increased the release of both (3H]NA and [14C]ACh in a concentration dependent manner. Atropine blocked the PDB-induced facilitation of ACh release but not the facilitation of NA release. 4. The protein kinase A (PKA) activator S Br-cMMP did not affect ACh release but enhanced NA release. 5. The PKC inhibitor H-7 (50-100 microM) inhibited the CS- or eserine-facilitated release evoked by IS. H 7 also inhibited 0.5 microM PDB-induced facilitation of ACh release but not NA release. 6. Down-regulating PKC by pretreatment for 30 min with 5 microM PDB decreased the facilitated release of ACh and the sessine-induced facilitation of NA release. 7. Electrically evoked contractions of the bladder strips exhibited a biphasic response to PDB (2.5 microM), which consisted of an initial enhancement of the peak amplitude and area followed after 20 min by an inhibition of contractions in a dose dependent fashion. 8. It is concluded that a phospholipase C-PKC signal transduction pathway is essential for muscarinic receptor-induced facilitation of ACh and NA release but is not involved in the non facilitated release of the path and NA release but is not involved in the non facilitated release of the post of the path and NA release but is not

L12 ANSWER 29 OF 121
ACCESSION NUMBER: 97
DOCUMENT NUMBER: PU
TITLE: Mu

AUTHOR: CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

EASTON NUMBER: 97077595 MEDLINE
UMENT NUMBER: 97077595 MEDLINE
UMENT NUMBER: Pubmed ID: 8920162
LE: Muscarinic receptor subtypes in the submandibular gland and the urinary bladder of the rabbit: in vivo and in vitro functional comparisons of receptor antagonists.

HOR: Tobin G Pepartment of Physiology and Pharmacology, Goteborg University, Sweden.

MECE: Journal of autonomic pharmacology, (1995 Dec) 15 (6) 451-63.

Journal of autonomic pharmacology, (1995 Dec) 15 (6) 451-63.

Journal code: 8106455. ISSN: 0144-1795.

LOUNTRY: ENGLAND: United Kingdom
UMENT TYPE: Journal; Article; (JOURNAL ARTICLE)
GUAGE: English
E SEGMENT: Priority Journals
RY MONTH: 199701
RY DATE: Entered STN: 19970128
Last Updated on STN: 19970128
Last Updated on STN: 19970128

Last Updated on STN: 1997018

1. In pentobarbitone anaesthetized rabbits, the inhibitory effects of muscarinic receptor antagonists with different selectivity profiles were examined on carbachol-evoked submandibular secretion and urinary bladder contractions, and on parasympathetically nerve-evoked secretion. On isolated submandibular studied on carbachol-evoked release of potassium and on the overflow of tritium in response to electrical field stimulation. 2. In vivo, 4-DAMP equipotently inhibited simultaneously carbachol-evoked submandibular secretory and contractile responses of the urinary bladder, while pirenepine was found to be four times as potent in inhibition of parasympathetic nerve evoked salivation. 2. In vivo, 4-DAMP submandibular secretory response compared with the contractile response. 3. The inhibition of carbachol-evoked submandibular inhibition of parasympathetic nerve evoked salivation.

Methoctramine secretory and contractive responses of the urinary bladder, while pirenepine was found to be four times as potent in inhibition of parasympathetic nerve evoked salivation.

Methoctramine secretory response compared with the contractile response of the urinary propose inhibition of parasympathetic nerve evoked salivation.

Methoctramine secretory secret

21 MEDLINE on STN
97060766 MEDLINE
PubMed ID: 8904812
Agents for the treatment of overactive detrusor. V.
Synthesis and inhibitory activity on detrusor contraction of N-tert-butyl-4,4-diphenyl-2-L12 ANSWER 31 OF 121
ACCESSION NUMBER: 9
DOCUMENT NUMBER: P
TITLE: A

cyclopentenylamine. Take K; Okumura K; Tsubaki K; Taniguchi K; Terai T; AUTHOR:

CORPORATE SOURCE:

Shiokawa Yesearch Laboratories, Fujisawa Pharmaceutical Co. Ltd., Osaka, Japan. Chemical & pharmaceutical bulletin, (1996 Oct) 44 (10) SOURCE:

1858-64; Journal code: 0377775, ISSN: 0009-2363.

Journal; Article; (JOURNAL ARTICLE)

English Priority Journals

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals

Y MONTH: 199612

Y DATE: Entered STN: 19970128

Last Updated on STN: 19970128

Last Updated on STN: 19970128

Entered Medline: 19961231

N-tert Butyl-4,4-diphenyl-2 cyclopentenylamine ((+/-) 3) was designed to restrict the conformation of terodline 1 and was synthesized in a 6 step approach starting with diphenylacetaldehyde (10) or in a 4-step approach starting with diphenylacetaldehyde (10) or in a 4-step approach starting with diphenylacetaldehyde (10) or in a 4-step approach starting with 2,2-diphenyl4-4-pentenoic acid (17). Using dip-tolucyltartaric acid as a resolving agent, the synthetic (+/-)-3 was resolved into its optically pure forms, (-)- and (+)-3. The (-)-enantiomer (-)-3.HCl (FKS44) showed about ten times more potent inhibitory activity on urinary bladder
rhythmic contraction in rats (ED30 = 0.18 mg/kg, i.v.) than terodiline (ED30 = 1.9 mg/kg, i.v.), while the (+)-enantiomer (+)-3.HCl showed no inhibitory activity at 1.0 mg/kg i.v. Compound (-)-3.HCl (FKS44) has pharmacological properties similar to those of terodiline, as evaluated by in vitro assay and is currently in clinical development for the treatment of overactive detrusor.

L12 ANSWER 32 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

MEDIANS ON SIN 97004322 MEDLINE PubMed ID: 8851633 Effect of thapsigargin on the contractile response of the normal and obstructed rabbit urinary TITLE:

bladder.

OR:
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OR:
ORDERORS SOURCE:
ORDERORS SOURCE:
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OPAPTE SOURCE:
OPAPTE SOURCE:
OPAPTE OF THE OFFICE OFFIC AUTHOR CORPORATE SOURCE:

CONTRACT NUMBER:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE

DR.Ro.1-33559 (NIDDK)
DR.Ro.1-44689 (NIDDK)

**RCE: Pharmacology, (1996 Feb) 52 (2) 119-24.

**Journal code: 0152016. ISSN: 0031-7012.

**COUNTRY: Switzerland
UMENT TYPE: Journal; Article; (JOURNAL ARTICLE)
GUAGE: English
E SEMPRIT: Priority Journals
RY MONTH: 199612

**RY MONTH: 19962

**RY MONTH: 19962

**Excitation-contraction coupling is achieved by translocation of calcium from the extracellular space as well as by the release of calcium from intracellular stores. Thapsigargin has been shown to selectively block the sarcoplasmic Ca-ATPase, thereby preventing the reuptake of calcium anto intracellular stores and the participation of these calcium storage sites in the contractile response to stimulation. The current study determined the effect of thapsigargin on the contractile response to field stimulation, bethanechol, and KCl in control rabbit bladders and bladders obtained from rabbits subjected to partial outlet obstruction. Partial bladder outlet obstruction resulted in a marked increase in bladder solated from control rabbits to field stimulation, bethanechol, or KCl. However, bladder strips isolated from control rabbits to field stimulation, obstructed rabbits showed a significant concentration dependent decrease in the contractile response to field stimulation in the presence of thapsigargin (5-40 microw) had no effect on the contractile response of bladder strips isolated from control rabbits to field stimulation, obstructed rabbits showed a significant concentration dependent decrease in the contractile response to field stimulation in the presence of thapsigargin. The response to field stimulation in the presence of thapsigargin. The response to field stimulation in the presence of thapsigargin for the sponse of bladder strips isolated from control rabbits to either bethanechol or KCl. In general, the data described in this study support our current hypothesis: as smooth muscle cells enlarge (hypertrophy) and the cell volume increases, there is an increased dependence on the release of intracellu

L12 ANSWER 34 OF 121 ACCESSION NUMBER: 9 MEDLINE on STN DOCUMENT NUMBER: TITLE:

CORPORATE SOURCE:

SOURCE:

21 MEDLINE on STN
95218053 MEDLINE
PubMed ID: 8641667
Toxicological comparison of a muscarinic agonist
given to rats by gavage or in the diet.
Dethloff LA: Chang T; Courtney C L
Department of Pathology, Parke-Davis Pharmaceutical
Research, Ann Arbor, MI 48105, USA.
Food and chemical toxicology: an international journal
published for the British Industrial Biological Research
Association, (1996 Apr) 34 (4) 407-22.
Journal code: 8207483. ISSN: 0278 6915.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: English ILE SEGMENT:

Priority Journals ENTRY MONTH ENTRY DATE:

English
ENGENT: Priority Journals
ENGENT: Entered STN: 19960726
Entered Medline: 19960717
Corneal opacities and urfnary tract sepsis were previously
observed by the authors in rats given muscarinic agonists mixed
in the diet or by gavage. To explain the differential toxicity generated
by each means of administration, toxicokinetics of the muscarinic
agonist CI:979 were investigated. In addition, the muscarinic
antagonist scopolamine was co-administered with CI-979 to evaluate the
relationship of these effects to pharmacological mechanism of action of
CI-979. Female rats were given CI-979 daily by gavage at 0, 1, 10 and 30
mg/kg body weight or in the diet at 0, 1, 10 and 50 mg/kg body weight for
up to 14 days. Dose-related clinical signs of muscarinic
stimulation, such as sialorrhoea and dacryorrhoea, were observed
predominantly in rats given 10 and 30 mg/kg body weight CI:979 by gavage,
and corresponded with the high plasma drug concentrations. In contrast,
hydronephrosis, pyelonephritis, and inflammation and necrosis of the
kidney, uninary bladder, urethra and urfnary
papilla were linked to sustained, albeit lower plasma drug concentrations
attained by dietary administration of CI-979 at 10 and 50 mg/kg body
weight. Comparable incidences of corneal opacities were induced by both
means of administration, but lesions appeared more rapidly and were
generally of greater severity when CI-979 was given in the diet. The
induction of corneal lesions, as well as urinary sepsie, may not
relate simply to maximum plasma concentrations or to areas under the curve
pear se, but rather may arise when plasma drug concentrations are
sustained. Corneal opacitication and development of urinary
tract pathology were inhibited by acopolamine, suggesting that
these effects were related to the muscarinic mechanism of action
of CI-979.

L12 ANSWER 33 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 96341615 MEDLINE
DOCUMENT NUMBER: PUMMed ID: 8750383
TITLE: Prodynamic and other effects of tolterodine: a novel antimuscarinic drug for the treatment of detrusor

SIGNAMESCATING GRUG FOR THE TREATMENT OF GETTUSOR
OVERACTIVITY
Stahl M M, Exetrom B; Sparf B; Mattiasson A; Andersson K E
Department of Clinical Pharmacology, Lund University
Hospital, Sweden.
Neurourology and urodynamics, (1995) 14 (6) 647-55.
Journal code: 8303326. ISSN: 0733 2467.
United States
Journal; Article; (JOURNAL ARTICLE)
Enolish AUTHOR: CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: English Priority Journals

E-SEMENT: Priority Journals
YY MONTH: 199610
YY MONTH: 199610
Tolterodine, a novel compound intended for treatment of urgency and urge incontinence, has been characterized as a potent muscarinic receptor antagonist in pharmacological in vitro and in vivo studies. In cats, tolerodine, as shown to reduce bladder pressure at doses significantly lower than those affecting salivation. To predict clinical effectiveness, an open pilot study was performed in healthy male volunteers. Efficacy was measured by cystometry and by spontaneously reported effects after administration of a single oral dose of tolterodine, 6.4 mg, given as a water solution. Tolterodine had distinct inhibitory effects on urinary bladder function, both at 1 and 5 hours post-dose. At 1 hour, but not at 5 hours post-dose tolterodine also significantly reduced stimulated salvation. In addition to the objectively demonstrated changes in urodynamic parameters, most volunteers experienced voiding difficulties. No significant changes in blood pressure, heart rate, or near point of accommodation were registered. Tolterodine, in the dosage used, was both objectively and subjectively shown to exert a marked inhibitory effect on micturition in healthy subjects, and the data suggest a more pronounced effect on bladder function than on salivation.

L12 ANSWER 35 OF 121 ACCESSION NUMBER: 90 DOCUMENT NUMBER: Pr 21 MEDLINE on STN 96162560 MEDLINE PubMed ID: 8583354

TITLE:

PubMed ID: 8583354
Analysis of the mechanisms underlying the contractile response induced by the hydroalcoholic extract of Phyllanthus urinaria in the guinea-pig urinary bladder in-vitro.
Dias M A; Campos A H; Cechinel Filho V; Yunes R A; Calixto

AUTHOR:

CORPORATE SOURCE:

J B Department of Pharmacology, Universidade Federal de Santa Catarina, Florianopolis SC, Brazil. Journal of pharmacy and pharmacology, (1995 Oct) 47 (10) 846-51 SOURCE:

846-51. Journal code: 0376363. ISSN: 0022-3573.

PUB. COUNTRY: DOCUMENT TYPE: ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE)

English

LANGUAGE: FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE:

SUAGE

SEMENT: Priority Journals
RY MONTH: 199603
RY DATE: Entered STN: 19960327
Last Updated on STN: 19980206
Entered Medline: 19960315
The hydroalcoholic extract of Phyllanthus urinaria (Euphorbiaceae), substance P and substance P methyl ester all caused graded contractions in the guinea-pig urinary bladder. Responses to hydroalcoholic extract and substance P were markedly inhibited in calcium-free Krebs solution, this effect being reversed by reintroduction of calcium in the medium. The contraction in response to hydroalcoholic extract and substance P were markedly inhibited in calcium-free Krebs solution, this effect being reversed by reintroduction of calcium in the medium. The contraction in response to hydroalcoholic extract was unaffected by atropine, propranolol, prazosin, yohimbine, tetrodotoxin, w conotoxin, nicardipine, HOE 140, guanethidine, staurosporine, phorbol ester or indomethacin, excluding the involvement of nervous mediated responses, or action via cholinergic, adrenergic, kinnins, cyclo-oxygenase metabolites, protein kinase C or activation of L or N-type calcium channels. The selective NK1 tachykinin antagonist (FK 888), but not NK2 (SR 48968) antagonized substance P-induced contraction, but both drugs failed to effect Phyllanthus urinaria-induced contraction. Prolonged desensitization of guinea pig urinary bladder with capsaicin (10 microm) or preincubation of guinea-pig urinary bladder with capsasepine did not affect contraction caused by hydroalcoholic extract. Ruthenium red almost completely abolished capsaicin-induced contraction, but had no effect on hydroalcoholic extract caused marked potentiation of the twitch response in the preparations field stimulated. The facilitatory effect of substance P, but not that of hydroalcoholic extract, was prevented by the NK1 (FK 888), but not by NK2 (SK 48968) antagonist. We concluded that contraction induced by hydroalcoholic extract of Phyllanthus urinaria in the guinea pig urinary bladder involves direct action on smooth muscle and relies on t

L12 ANSWER 36 OF 121 ACCESSION NUMBER: 96 MEDLINE on STN DOCUMENT NUMBER:

AUTHOR CORPORATE SOURCE:

All MEDLINE On STN
96160127 MEDLINE
PubMed ID: 8588301
A review of nonpesticide phosphate ester-induced
neurotoxicity in cattle.
Coppock R W; Mostrom M S; Khan A A; Stair E L
Environmental Toxicology Research, Alberta Environmental
Centre, Vegreville, Canada.
Veterinary and human toxicology, (1995 Dec) 37 (6) 576-9.
Ref: 20 SOURCE

Ref: 20 Journal code: 7704194. ISSN: 0145-6296.

PUB. COUNTRY: DOCUMENT TYPE:

Journal code: 7704194. ISSN: 0145-6 United States Journal; Article: (JOURNAL ARTICLE) General Review; (REVIEW) (REVIEW, TUTORIAL) English Friority Journals

LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

SEMENT: Friority Journals

WY MONTH: 199603

WY MONTH: 19960404

Last Updated on STM: 19960404

Entered Medline: 19960327

Nonpesticide phosphate esters induce delayed neurotoxicity in cattle. The most common exposures are to complex mixtures of triaryl phosphate used in lubricating oils. Oral ingestion is most common, but dermal exposures have also occurred. Clinical signs of Cholinesterase (ChE)

inhibition may or may not be seen. Depending on the biochemical targets, the percent reduction in blood ChE is variable and can be < 30% of normal activity. Organophosphate ester-induced delayed neurotoxicity cannot be predicted by inhibition of blood ChEs. Signs of delayed neurotoxicity occur 2 to 25 d after exposure; these signs are neurologic deficiencies of the antigravity muscles and the muscles of the urinary bladder and larynx. Affected cattle may dribble urine and some may be mute. Signs of ChE inhibition generally are not observed in animals with neurological deficiencies. Pathologic findigs are axonopathy and myelin degeneration of nerves with long axons located in both the peripheral and central nervous systems. In the spinal coord, location of the affected nerve tracts is variable. Degenerative changes occur in motor neurons. Calves are less susceptible to organophosphate ester-induced delayed neurotoxicity than cows. A dose of 500 mg triaryl phosphate/kg body weight will produce complete paralysis in a mature cow in 26 d.

L12 ANSWER 38 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: P MEDLINE on STN 96022482 MEDLINE PubMed ID: 7580750

Pubmed 10: 7580750 [The role of protein kinase C and muscarinic cholinoreceptors in vasopressin-stimulated water

transport]. Or oli proteinkinazy C i muskarinovykh kholinoretseptorov v stimulirovannom vazopressinom vodnom transporte.
Bagrov Ia Iu; Dmitrieva N I, Manusova N B
Eksperimental'naia i klinicheskaia farmakologiia, (1995
Jul-Aug) 58 (4) 33-5.
Journal code: 9215981. ISSN: 0869 2092.
RUSSTA: Russian Federation
Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

Russian Priority Journals FILE SEGMENT:

AUTHOR:

ENTRY MONTH: ENTRY DATE:

RY MONTH: 199512

Entered STN: 19960124

Last Updated on STN: 19960124

Entered Medline: 19951213

The role of protein kinase C (PKC) in the control of vasopressinstimulated water transport in the frog urinary bladder and its modulation by M2:agonist oxotremorine has been studied. Using the PKC inhibitor, staurosporine we showed that PKC in the region pf the basal membrane suppressed vasopressin-stimulated water transport, whereas PKC in the apical region potentiated this transport. It was also found that from the two types of oxotremorine action on stimulated water transport determined by its concentration only inhibition is mediated through PKC.

L12 ANSWER 37 OF 121
ACCESSION NUMBER: 9
DOCUMENT NUMBER: P
TITLE: ME MEDLINE on STN 96073411 MEDLINE PubMed ID: 7576597

Ruscarinic suppression of Ca2+ current in smooth muscle cells of the guinea-pig urinary bladder.

AUTHOR: CORPORATE SOURCE:

bladder.
Yoshino M; Yabu H
Department of Physiology, Sapporo Medical University, Japan. Specimental physiology, (1995 Jul) 80 (4) 575-87. Journal code: 9002940. ISSN: 0958-0670. BKGLAND: United Kingdom Journal; Article: (JOURNAL ARTICLE)

English

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: Priority Journals 199512

GUAGE:

English

E SEGMENT: Priority Journals

RY MONTH: 199512

RY DATE: Entered STN: 19960124

Last Updated on STN: 20000303

Entered Medline: 19951205

The suppressive action of carbachol (CCh) on the Ca2+ current (ICa) in smooth muscle cells of the guinea pig urinary bladder

was investigated using the whole-cell patch clamp technique. Bath application of 10 microM CCh reduced the amplitude of ICa by 92 */- 3.8% (n = 9). Adding 1 microM atropine to the bath completely blocked the action of CCh indicating that the suppressive action of CCh on ICa is mediated by the activation of muscarinic receptors.

Intracellular perfusion of the mon-hydrolysable GTP analogue, guanosine 5'-0-(3-thiotriphosphate) (GTP gamma S; 200 microM) mimicked the effects of CCh. Sustained suppression of ICa was observed when GTP gamma S was present in the cytoplasm. Intracellular perfusion of inositol 1,4,5-trisphosphate (InsP 3; 20 microM) also suppressed ICa: its effect was not sustained but transient. The protein kinase C activator, phorbol ICa. When intracellular Ca2+ was strongly buffered by the Ca2+ chelator SOTA (20 MM) in the patch pipette, the sustained suppression of ICa was abolished. Inclusion of 3 mg/ml hepartn, a blocker of InsP3-induced Ca2+ release, in the patch pipette reduced the degree of sustained ICa suppression by 43.2 */- 1.9% (n = 7). Adding thapsigargin (TG), a sarcoplasmic reticulum Ca2+-Arase inhibitor, to a wash solution reduced the recovery of ICa by about 50% suggesting that approximately half of the ICa suppression internal Ca2+ stores. From these results it appears that CCh suppressive internal Ca2+ chelase from TO-sensitive internal Ca2+ chelase. From these results it appears that CCh suppress ICa vis two independent mechanisms. (1) Ca(2+) mediated inactivation of the Ca2+ chelanel, which is caused by C2+ release from InsP3- and TO sensitive internal stores, and (2) a GTP-binding protein-mediated mechanism, which requires intracellular Ca2+.

L12 ANSWER 39 OF 121
ACCESSION NUMBER: 9
DOCUMENT NUMBER: P
TITLE: I 21 MEDLINE on STN 96007972 MEDLINE PubMed ID: 8566107 In vivo and in vitro effects of muscarinic

receptor antagonists on contractions and release of [3H]acetylcholine in the rabbit urinary

AUTHOR: CORPORATE SOURCE:

SOURCE:

bladder. Tobin G; Sjogren C Department of Pharmacology, University of Goteborg, Sweden. European journal of pharmacology, (1995 Jul 25) 281 (1)

Journal code: 1254354. ISSN: 0014-2999. Netherlands

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

Journal; Article; (JOURNAL ARTICLE) English

Priority Journals

L12 ANSWER 40 OF 121 1 MEDLINE on STN 95203909 MEDLINE ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

21 MEDLINE ON STM
95203909 MEDLINE
PubMed ID: 7896306
Effects of adenosine on contractility and 45Ca-uptake in
rat urinary bladder.
Farija S C; Raviprakash V; Mishra S K
Division of Pharmacology and Toxicology, Indian Veterinary
Research Institute, Izatnagar.
Indian journal of experimental biology, (1994 Nov) 32 (11)
781-5. CORPORATE SOURCE:

SOURCE:

Journal code: 0233411. ISSN: 0019-5189. PUB. COUNTRY: India

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE: LANGUAGE:

English Priority Journals 199504 FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

5 SEGNENT: Priority Journals
YM MONTH: 199504
YM MONTH: 199504
Entered STN: 19950504
Last Updated on STN: 19950504
Entered Medline: 19950425
Effects of adenosine on K+ and ACh-stimulated contractility and 45Ca uptake were studied in the rat urinary bladder smooth muscle and were compared with those of nifedipine. Both adenosine (10(-5) M) and nifedipine (10(-7) M/10(-8) M) significantly inhibited the contractions elicited by K+ (10(2)-21x 10(-2) M), Ca2+ (10(-4)-3 x 10(-2 M) in K(+) depolarized preparations and ACh (10(-9) M-3 x 10(-3) M). Further, adenosine (10(-5) M) significantly (F< 0.05) inhibited K+ (10(-1) M)-stimulated 45Ca uptake in the bladder strips.
However, it had little effect on inward 45Ca movement resulting from ACh (10(-4) M)-induced stimulation. On the other hand, nifedipine (10(-7) M) significantly (F< 0.001) reduced both K+ and ACh-induced 45Ca-uptake in this tissue. It is concluded that the calcium channels activated by addenosine is limited to C2+ uptake through voltage operated calcium channels, while receptor operated calcium channels activated by muscarinic receptor stimulation appear to be insensitive to the

ANSWER 42 OF 121 MEDLINE on STN 95151079 MEDLINE PubMed ID: 7848339 ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

AUTHOR:

PubMed ID: 7949379

Wrinary bladder-selective action of the new antimuscarinic compound vamicamide. Oyasu H; Yamamoto T; Sato N; Sawada T; Ozaki R; Mukai T; Ozaki T; Nishii M; Sato H; Fujiwara T; + Pharmacological Research Laboratories, Fujisawa Pharmaceutical Co., Ltd., Osaka, Japan. Arzneimittel-Forschung, (1994 Nov) 44 (11) 1242-9. Journal code: 0372650 ISSN: 0004-4172. CERNANY; Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE: English

Priority Journals

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SAGENT: English

5 SEGMENT: Priority Journals

19503

17 MONTH: 199503

18 DATE: Entered STN: 19950316

Entered Medline: 19950306

1. The inhibitory action of vamicamide (FK176, ('/-)-(2R*,4R*)-4-d-methylamino-2-phenyl-2-(2-pyridyl)valeramide, CAS
132373-81-0) on the responses of various tissues to the cholinergic agonists, carbachol and McN-A-343 (4-(m-chlorophenylcarbamoyloxyl-2-butynyl-trimethylammonium chloride, CAS 55-45 8), was investigated in isolated tissue preparations. Vamicamide showed competitive antagonistic actions against all the preparations tested and its pA2 value for the urinary bladdar was 6.82, which was higher than that for the atria (5.94) and almost the same as that for the vas deferens (6.90) and for the atomach (6.81). The pA2 values of oxybutynin hydrochloride (oxybutynin) and atropine sulfate monohydrate (atropine) were nearly the same in all the tissues tested 2. Oral administration of vamicamide 0.1-1.0 mg/kg inhibited dose-dependently spontaneous bladdar contractions caused by raising the intravesical volume in conscious rats. Inhibitory actions were also obtained with 0.32-3.2 mg/kg of oxybutynin was shorter than that of vamicamide or atropine. Vamicamide further inhibited bladdar contractions in rats following intravesical administration of 0.05-0.5 mg/kg ml solution. 3. Vamicamide had no effect or only slightly inhibited spontaneous motility of the stomach and distal colon in conscious rats, as well as heart rate and salivary secretion in conscious doss, after oral dosing with 3.2 mg/kg of the compound. Similar results were obtained with oxybutynin, excepting the occurrence of tachycardia. (ABSTRACT TRUNCATED AT 250 WORDS)

L12 ANSWER 41 OF 121 MEDLINE on STN
ACCESSION NUMBER: 95156307 MEDLINE
DOCUMENT NUMBER: 9516407 MEDLINE
TITLE: Minuscarinic receptor-mediated facilitation of acetylcholine release in the rat urinary bladder.

AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

LE: MI muscarinic receptor-mediated facilitation of acetylcholine release in the rat urinary bladder.

HOR: Somogyi G T; Tanowitz M; de Groat W C
Department of Pharmacology, University of Pittsburgh, PA 15261.

TRACT NUMBER: MH 30915 (NIMH)
Journal of physiology, (1994 Oct 1) 480 (Pt 1) 81-9.
Journal code: 0265262. ISSN: 0022-3751.

ENGLAND: United Kingdom
JOURNATY: MOUTH JOURNAL ARTICLE)
SUAGE: ENGLAND: United Kingdom
JOURNAT TYPE: JOURNAL ARTICLE)
SUAGE: English
EN HONTH: Priority Journals
RY DATE: Entered STN: 19950322
Last Updated on STN: 19950322
Last Updated on STN: 19950310

1. Release of [3H]ACh in response to electrical field stimulation (10 Hz) was measured in strips of rat urinary bladder and cardiac atrial tissues previously incubated with [3H]choline. 2. The volley output of [3H]ACh release was positively correlated with frequency of stimulation in the urinary bladder but negatively correlated in the arrium. 3. The quantity of [3H]ACh release was influenced by the pattern and duration of stimulation. Continuous stimulation (CS) with trains of 100 shocks released 10 times larger amounts of ACh than the same number of shocks presented as short trains of intermittent stimulation (TS): ten shocks per train with 5 s inter-train intervals. 4. The facilitation of transmitter release was nother trains agent, markedly facilitated ACh release induced by CS and IS. This effect was blocked by atropine. 5. Release of ACh from atrial strips did not exhibit CS induced facilitation. Eserine decreased IS- and CS-evoked ACh release in the atrium. 6. It is concluded that continuous stimulation of postganglionic cholinergic nerves in the rat urinary bladder leads to the activation of M1 muscarinic, facilitatiory presynaptic receptors which enhance the release of ACh. Presynaptic facilitation may be an important mechanism for modulating neural input to the bladder during micturition.

L12 ANSWER 43 OF 121 MEDLINE on STN
ACCESSION NUMBER: 95147990 MEDLINE
DOCUMENT NUMBER: 95147990 MEDLINE
TITLE: pubMed ID: 7845476
The role of extracellular Ca2+ in carbachol induced tonic concraction of the pig detrusor smooth muscle.

AUTHOR: Uchida W, Masuda N, Shirai Y, Shibasaki K, Satoh N;
Takenada T
CORPORATE SOURCE: Cardiovascular and Atherosclerosis Research Laboratories,
Yamanouchi Institute for Drug Discovery Research, Ibaraki,
Japan.

Japan: Naunyn-Schmiedeberg's archives of pharm 350 (4) 398-402.
Journal code: 0326264. ISSN: 0028-1298. GERMANY: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) SOURCE: -Schmiedeberg's archives of pharmacology, (1994 Oct)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

. COUNTRY: GERMAN: GERMANY: GERMANY: GERMANY: MEMORY TYPE: Journal; Article; (JOURNAL ARTICLE)

JUACE: English
E SEGMENT: Priority Journals

RY MONTH: 199503

RY DATE: Entered STN: 19950316

Last Updated on STN: 19950316

Entered Medline: 19950307

The role of extracellular Ca2+ in the tonic-contractile response to muscarinic receptor stimularion was investigated in isolated detrusor smooth muscle from the pig urinary bladder.

Carbachol (10(8):10(-5) M) produced a concentration-dependent contractile response in isolated pig detrusor smooth muscle strips consisting of an initial phasic component followed by a tonic component. During the plateau of the tonic contractions induced by carbachol at the submaximal concentration of 10(-6) M, the inhibiting effects of atropine, EGTA, nitedipine (a voltage-dependent calcium channel antagonist), H-7 [a protein kinase C (FCC) inhibitor] and YM934 (a pocassium channel opener) on the contractions were evaluated. Atropine (10(-10)-1 x 10(-8) M) concentration-dependently inhibited the tonic contractions induced by carbachol. In the same experimental conditions, EGTA (4 mM) and nifedipine (10(-9)-1 x 10(7) M) depressed the tonic contractions in a concentration-dependent manner as did H-7 (10(-5)-2 x 10(-5) M) and YM934 (10(-6) M). Nowever, H-7 (10(-5)-3 x 10(-5) M) and YM934 (10(-6) M) were very weak in inhibiting the contractions induced by KCl (50 mM) in isolated pig detrusor smooth muscle strips. These results suggest that the tonic-contractile response induced by carbachol in pig detrusor smooth muscle strips is dependent mainly on depolarization of the cell membranes and an influx of extracellular Ca2+, and also suggest that this depolarizing response may be due to inactivation of ATP-sensitive potassium channels through muscerinic activation of PKC.

ANSWER 44 OF 121 11 MEDLINE ON STN 94287486 MEDLINE PubMed ID: 8016895 Muscarinic acetylcholine receptor subtypes in ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

smooth muscle.

COMMENT:

Smooth muscle.

Comment on: Ugeskr Laeger. 1993 Aug 9;155(32):2438-42.

PubMed ID: 8102824

Comment in: Trends Pharmacol Sci. 1994 Nov;15(11):407-8.

PubMed ID: 7855902

Eglen R M; Reddy H; Watson N; Challiss R A
Institute of Pharmacology, Syntex Discovery Research, Palo
Alto, CA 94304.

Trends in pharmacological sciences, (1994 Apr) 15 (4)

114-9. Ref: 50

Journal code: 7906158. ISSN: 0165 6147.

ENGLAND: United Kingdom

Commentary

CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

ENGLAND: United Kingdom
Commentary
Journal, Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
(REVIEW, ACADEMIC)
English
Priority Journals
199407

LANGUAGE: FILE SEGMENT:

AUTHOR:

ENTRY MONTH:

ENTRY DATE

SEGMENT: Priority Journals
IY MONTH: 199407
IY DATE: Entered STN: 19940810
Last Updated on STN: 20000303
Entered Molline: 19940728
Muscarinic acetylcholine M2 and M3 receptor subtypes are
coexpressed in many types of smooth muscle including gastrointestinal
smooth muscle, urinary bladdar and vascular and airway
tissue. Activation of M3 receptors, via the G protein Gq, results in
increased polyphosphoinositide hydrolysis, release of Ca2+ ions from the
sarcoplasmic reticulum and consequently causes contraction. Quantitation
of the relative expression of M2 and M3 receptors has shown that the
proportion of M2 receptors often predominates over the M3 receptor
population by 4:10 rmore. Although it is established that M2 receptors
preferentially link, via a pertussis toxin-sensitive G protein Gi, to
inhibition of adenylate cyclase activity, relatively little is
known concerning the physiological role of the M2 receptor population. In
this review, Richard Bylen and colleagues discuss recent data concerning
the possible role(s) of muscarinic receptor subtypes in smooth
muscle and appraise the pharmacological methods for dissecting the
function of muscarinic receptor subtypes in tissues
Co expressing multiple receptors.

ANSWER 46 OF 121 MEDLINE on STN L12 ANSWER 46 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

21 MEDLINE on STN 94106595 MEDLINE PubMed ID: 8279533 Muscarinic inhibition of ATP-sensitive

CORPORATE SOURCE:

CONTRACT NUMBER:

Muscarinic inhibition of ATP-sensitive
K+ channels by protein kinase C in urinary
bladder smooth muscle.
Bonev A D; Nelson M T
Department of Pharmacology, University of Vermont,
Colchester 05446-2500.
HL-44455 (NHLBI)
American journal of physiology, (1993 Dec) 265 (6 Pt 1)
C1723-8.
Journal code: 0370511. ISSN: 0002-9513.

PUB. COUNTRY: United States Journal; Article; (JOURNAL ARTICLE)

English

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: Priority Journals 199402

E SEOMENT: Priority Journals
RY MOXTH: 1994021
RY DATE: Shtered STN: 19940218
Last Updated on STN: 20021219
Entered Medline: 19940208
We explored the possibility that muscarinic receptor stimulation
can inhibit ATP-sensitive K+ (KATP) channels in smooth muscle
cells from guinea pig urinary bladder. Whole cell K+
currents were measured in smooth muscle cells isolated from the detrusor
muscle of the guinea pig bladder. Stimulation of
muscarinic receptors by carbachol (CCh; 10 microM)
inhibited KATP currents by 60.7%. Guanosine 5'-O- (2thiodiphosphate) in the pipette (internal) solution prevented the
CCh-induced inhibition of KATP currents. Activators of protein
kinase C (PKC), a diacylglycerol analogue, and phorbol 12-myristate
13-acetate inhibited KATP currents by 63.5 and 73.9%,
respectively. Blockers of PKC (bisindolylmaleimide GF-109203X and
calphostin C) greatly reduced CCh inhibition of KATP currents.

KATP channels in smooth muscle cells from urinary
bladder through activation of PKC.

L12 ANSWER 47 OF 121
ACCESSION NUMBER: 94
DOCUMENT NUMBER: POTITLE: Ef

L12 ANSWER 45 OF 121 MEDLINE ON STN
ACCESSION NUMBER:

PUBMED ID: \$133908

Inhibitory effects of propiverine on rat and
guinea-pig urinary bladdar muscle.

AUTHOR:

COMPORATE SOURCE:

SOURCE:

Namyn-Schmiedeberg's archives of pharmacology, (1993 Dec)
348 (6) 559 62.

JOURNAL TYPE:

JOURNAL ARTICLE)

LANGUAGE:

ENTRY DATE:

ENTRY DATE:

Entered STN: 19940428

Entered Medline: 19940428

Last Updated on STN: 19940428

Entered Medline: 19940428

Last Updated on STN: 19940428

In muscle strips isolated from guinea-pig and rat urinary
bladder, propiverine (3-10 microw) inhibited

carbachol-induced contractions in the presence of verapamil and
Ca(2+)-induced contractions in access K+ medium containing atropine,
suggesting it has both anticholinergic and Ca(2+ channel blocking action was also demonstrated by recording inward
Ca(2+ currents in single cells dispersed from both species. The
ca(2+ channel blocking action was also demonstrated by recording inward
Ca(2+ currents in single cells dispersed from both species. The
inhibition of inward currents by propiverine was three times
stronger in the rat than the guinea-pig, IDSO being 7 microm for rat and
21 microm for guinea-pig. The recovery of the current after washout was
faster than that of mechanical inhibition. It is concluded that
propiverine blocks not only muscarinic receptors, but also Ca(2+
channels at similar concentrations.

current in smooth muscle cells isolated from guinea-pig urinary bladder.

Nakayama S University Department of Pharmacology, Oxford.

CE: British journal of pharmacology, (1993 Sep) 110 (1) 317 25.

Journal code: 7502536. ISSN: 0007-1188.

COUNTRY: ENGLAND: United Kingdom
JOURNAL ARTICLE)

SMARE: English:

ENGLAND: United Kingdom
JOURNAL ARTICLE)

SMARE: English:

ENGLAND: United Kingdom
JOURNAL ARTICLE)

SMARE: Priority Journals

ENGLAND: United Kingdom

YM MONTH: 199312

1. A whole-cell voltage clamp technique was used to examine the effects of purinoceptor and muscarinio receptor agonists on voltage-ensitive Cel* channels in guinea-pig isolated urinary bladder cells. 2. When the cell membrane was clamped at the holding potential, rapid application of ATP elicited a large inward current in normal solution containing 2.5 mM Car*, and reduced the subsequent Ca2+ channel current evoked by a depolarizing pulse (0 mV). Carbachol (CCh) elicited little membrane current, but similarly reduced the Ca2+ current. 3. When purinoceptor agonists were rapidly applied during conditioning depolarizations at +80 mV, an outward current was elicited, and the Ca2+ channel current evoked by the subsequent test potential of 0 mV was not affected. Application of CCh at +80 mV also elicited an outward current, but it reduced the subsequent test potential of 0 mV was not affected. Application of CCh at +80 mV also elicited an outward current was elicited and the Ca2+ channels current was attenuated by caffeine (10 mM).

5. In Ca(2+)-free, low Mg2+ solution, a Na+ current flowing through voltage-dependent Ca2+ channels was evoked by depolarization. This current was not reduced by bath application of purinoceptor agonists (ATP and alpha, beta-methylene ATP). 6. These results suggest that the main effect of purinoceptor stimulation is opening of non-selective cairon channels, and that **muscarinic** estimulation triggers Ca2+ channels are innac AUTHOR: CORPORATE SOURCE: SOURCE: PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

MEDLINE on STN
94035894 MEDLINE
PubMed ID: 8106107
Effects of excitatory neurotransmitters on Ca2+ channel current in smooth muscle cells isolated from guinea-pig wrinary bladder.

L12 ANSWER 48 OF 121 ACCESSION NUMBER: 93 DOCUMENT NUMBER:

AUTHOR CORPORATE SOURCE:

21 MEDLINE on STN
93374286 MEDLINE
PubMed ID: 8365654
Role of calcium in mediating the biphasic contraction of
the rabbit urinary bladder.
Zhao Y; Wein A J; Levin R M
Division of Urology, University of Pennsylvania School of
Medicine, Philadelphia 19104.
P50-DK-99257 (NIDDK)

CONTRACT NUMBER:

RO-1-DK 26508 (NIDDK) RO-1-DK133559 (NIDDK)

General pharmacology, (1993 May) 24 (3) 727-31. Journal code: 7602417. ISSN: 0306-3623. ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE:

LANGUAGE: English Priority Journals

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SIGNAMEN: Biggism : Biggism : Signamen : Sig

ANSWER 50 OF 121 1 MEDLINE on STN 93247153 MEDLINE PubMed ID: 8387115 L12 ANSWER 50 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

PubMed ID: 8387115
The pathophysiology of contractile activity in the chronic decentralized feline bladder.
Skehan A M; Downie J W; Awad S A
Department of Urology, Dalhousie University, Halifax, Nova Scotia, Canada.
Journal of urology, (1993 May) 149 (5) 1156-64.
Journal code: 0376374. ISSN: 0022-5347.
United States
Journal; Article; (JOURNAL ARTICLE)
English

CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

English

LANGUAGE: FILE SEGMENT: ENTRY MONTH: Abridged Index Medicus Journals; Priority Journals

ENTRY DATE:

SEGMENT: Abridged Index Medicus Journals; Priority Journals IT MONTH: 19305

XT MONTH: 19305

XT MONTH: Entered STN: 1930618

Last Updated on STN: 1930618

Entered Medline: 19930528

Autonomous wave activity occurs in the decentralized bladder and may contribute to upper tract damage and incontinence. In order to clarify the poorly understood pathophysiology and neuropharmacology of autonomous waves, cats were prepared with LT-S3 ventral rhizotomy alone or with LT-S3 ventral rhizotomy with and without total sympathectomy. The incidence of autonomous waves was < 15% 12 weeks after ventral or ventradorsal rhizotomy, but acute sympathectomy at 13 weeks increased the incidence to 58% in these groups. With chronic sympathectomy the incidence was 100%. This suggests that the waves arise locally via a mechanism which is independent of LT S3 dorsal rocts, due to lack of a suppressive sympathetic pathway. Autonomous waves were inhibited by arropine after acute sympathectomy and by prazosin after chronic sympathectomy, but increased inhibition occurred after both drugs in either case. Adrenergic neuron depletion with 6-hydroxydopamine enhanced wave activity, which was incompletely inhibited by subsequent atropine. This implies that the peripheral reflex pathway has facilitatory alpha 1-adrenergic, muscarinic and also noncholinergic nonadrenergic elements.

Clinically, sensory or sympathetic damage caused incontinence, but sympathectomy also caused high pressure waves, which may cause upper tract damage and treatment resistant incontinence in patients.

ANSWER 49 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

SOUTHOR

21 MEDLINE On STN
93251320 MEDLINE
PubMed ID: 8485522
Differential changes of adrenoceptor- and
muscarinic receptor-linked prostacyclin synthesis
by the acrts and wrinary bladder of the
diabetic rr.
Jeremy J Y; Thompson C S; Mikhailidis D P
Department of Chemical Pathology and Human Metabolism,
Royal Free Hospital and School of Medicine, University of
London. CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

Department of Chemical Pathology and Human Metabolism, Royal Free Hospital and School of Medicine, University of London.

RCE: British journal of pharmacology, (1993 Apr) 108 (4) 1131-6. Journal code: 7502516, ISSN: 0007-1188.

COUNTRY: ENGLAND: United Kingdom University of Journal Code: 7502516, ISSN: 0007-1188.

COUNTRY: ENGLAND: United Kingdom University of Journal; Article; (JOURNAL ARTICLE) SIGNOE: English ESEGMENT: Priority Journals RY MONTH: 199366

RY MONTH: 199366

RY DATE: Entered STN: 19930618

Last Updated on STN: 19930604

1. The effect of experimental diabetes mellitus (DM; hyperglycaemic, non-ketototic; 2 months duration) in the rat on receptor-linked prostacyclin (PGI2) synthesis (measured as 6-cox-0-0971 alpha by radioimmunoassay) was studied in the acrts and wrinary bladder using adrenaline, anjotensin II (AII) and accylcholine (ACh). Signal transduction systems were studied via stimulation of PGI2 synthesis with phorbol ester dibutyrate (PBU), a protein kinase cativator (PKC), Ca2+ ionophore A23187 (A23187) and thapsigargin (both elevate intracellular Ca2+, activating phospholipase A2 (PLA2)) and arachidonate (AA) substrate for PGI2 synthesis 2. In response to adrenaline, AII and phorbol ester, active PGI2 release was markedly reduced (all > 754) in diabetic rats compared to controls. EC50s of the dose-response curves for adrenaline, AII and PDBU were also markedly increased in acrtse from DM rats compared to controls. Although there was decreased output of PGI2 in response to A23187 by acrtse from diabetic rats compared to controls 2 //-0.18 x 10(-6) M). There were no differences in PGI2 release for in the EC50s (mean +/-6.e. marked) increase in RGI2 output in response to A24187 and a marked decrease in EC50s for the ACh-PGI2 dose-response curves and diabetic rats (EC50 = 5.8 +/-0.18 x 10(-6) M). (ABSTRACT TRUNCATED AT 250 WORDS)

L12 ANSWER 51 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: P

TITLE:

AUTHOR:

CORPORATE SOURCE: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

EASION NUMBER: 93207573 MEDILINE
UMENT NUMBER: Pubmed 1D: 838453
LE: Effects of terflavoxate on stimulated contractions of urinary bladder in vitro.

HOR: Testa R, Guarneri L, Bernasconi P, Angelico P; Ibba M;
POGRATE SOURCE: Pharmacology Laboratories, Recordati S.p.A., Milan, Italy.

POGRATE SOURCE: Pharmacology Laboratories, Recordati S.p.A., Milan, Italy.

Arrneimittel-Forschung, (1993 Feb) 43 (2) 122-8.

JOURNAY: GERMANY: Germany, Federal Republic of
UMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

GUAGE: English

S SEDMENT: Priority Journals

RY MONTH: 199304

RY DATE: Entered STN: 19930507

Last Updated on STN: 19930507

Last Updated on STN: 19930507

Last Updated on STN: 19930507

The antispassmotic activity of terflavoxate (CAS 86433-39-8), a flavone derivative with spassmolytic properties on the urinary tract, has been studied in vitro, in comparison to the most common drugs utilized in the therapy of overactive detrusor, namely flavoxate, oxybutynin, and terodiline. Terflavoxate showed affinity for bladder (and brain) muscarinic receptors at micromolar level, however, its activity on carbachol-induced contractions of rat bladder was clearly non competitive, indicating that the compound is devoid of functional antimuscarinic property. Moreover, the observation that unlike antimuscarinic drugs, terflavoxate inhibited by more than 504 field stimulation-induced contractions of rabbit bladder strips, indicates that mechanisms other than the anticholinergic one should be responsible for its smooth muscle relaxant properties. Terflavoxate, flavoxate, oxybutynin, and terodiline were equally effective in inhibiting the two components of K(+)-induced contractions. In addition, while nifedipine and nicardipine were equally effective in inhibiting tonic than phasic contractions. In addition, while nifedipine and nicardipine were equally effective in manner calcium-induced contractions of potassium-depolarized bladder strips, the other spasmolytics behaved as mixed antagonists. Differences in calcium antagonistic

L12 ANSWER 52 OF 121 ACCESSION NUMBER: 9: DOCUMENT NUMBER: TITLE:

21 MEDLINE on STN
93112409 MEDLINE
PubMed ID: 1282072
A pharmacological study of NK1 and NK2 tachykinin receptor characteristics in the rat isolated urinary

AUTHOR: CORPORATE SOURCE:

SOURCE:

Characteristics in the lat sevents — bladder.

Hall J M; Flowers J M; Morton I K
Pharmacology Group, King's College London, London,
British journal of pharmacology, (1992 Nov) 107 (3) 777-84.

Journal code: 7502536. ISSN: 0007-1188.

ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

Enolish PUB. COUNTRY:

DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT: Priority Journals ENTRY DATE:

English

S SERMENT: Priority Journals

YY MONTH: 199302

Last Updated on STN: 19930219

Last Updated on STN: 19930202

1. We have estimated potencies of tachykinin receptor agonist and antagonist analogues in order to determine the recognition characteristics of tachykinin receptors mediating phasic contractile responses of the rat isolated urinary bladdar in vitro. 2. The

NK1-selective synthetic agonists, substance P methyl ester and GR73612, the synthetic NK2-selective agonists [beca-Ala8]-NKA(4-10) and GR64349, and the mammalian tachykinins, neurokinin A and neurokinin B, were assayed relative to substance P and were found to be approximately equipotent.

The NK3-selective agonist, senktide, was inactive (10 microW). 3.

Dotencies of all these agonists were not significantly different (P > 0.05) when experiments were carried out in the presence of the neutral endopeptidase inhibitor, phosphoramidon, and the kininase II inhibitor, enalagrilat (both 1 microW). 4. The NK1-selective antagonist, GR82334, inhibited responses to substance P methyl ester in a competitive manner in the rat urinary bladdar substance P methyl ester in a competitive manner in the rat urinary bladdar (5.38) and rat ileum (6.56) compared to the guinea-pig lleum (7.42). GR82334 (3 microW) was inactive against responses of the rat bladdar (6.38) and rat ileum (6.56) compared to the guinea-pig lleum (7.42). GR82334 (3 microW) was inactive against responses of the rat bladdar (6.38) and rat ileum to substance P methyl ester; however, in the rat bladdar at 1 microW, this antagonist reversibly inhibited responses both to the NK2 selective agonist (beta Ala8) NKA(4-10) and to the muscariaic agonist carbachol (P < or - 0.01), thus showing evidence of some non-selective depressant actions (ABSTRACT TRUNCATED AT 250 WORDS)

L12 ANSWER 54 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 93063387 MEDLINE
DOCUMENT NUMBER: Pubmed 1D: 1436123
TITLE: Pubmed 1D: 1436123
TITLE: Substance agonist and antagonist selectivity at muscarinic receptors in guinea-pig smooth muscles

AUTHOR:

CORPORATE SOURCE:

SOURCE:

muscarinic receptors in guinea-pig smooth muscles and cardiac atria.

Dorofeeva N A; Shelkovnikov S A; Starshinova L A; Danilov A F; Nedoma J; Tucek S
Sechenov Institute of Evolutionary Physiology and Biochemistry, Russian Academy of Sciences, St. Petersburg.
Naunyn Schmiedeberg's archives of pharmacology, (1992 Oct)
346 (4) 383-90.

Journal code: 0326264, ISSN: 0028-1298.

GERMANY; Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE) PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

English FILE SEGMENT: ENTRY MONTH: Priority Journals

ENTRY DATE:

SEGMENT: Priority Journals

STEGMENT: Priority Journals

YMONTH: 199212

Last Updated on STN: 19930122

Entered Medline: 19921201

Fotencies of 11 muscarinic agonists in eliciting contraction of smooth muscle in quintea-pig ileum, trachea, urinary

bladder and uterus and in inhibiting the rate of contractions of cardiac atria were compared. While acetylcholine (ACh) was the most potent agonist on the ileum, uterus and cardiac atria, cis-L(+)-dioxolane was equally as potent as ACh on the ileum and more potent on the urinary bladder and trachea. Compared

to ACh, methyl furnethide, oxortemorine, acetoxybut-2-inyl-trimethylammonium and cis-L(+)-dioxolane acted weakly on the atria. Aceclidine, arecoline and acetyl-beta-methyl-tohline displayed selectivity for the urinary bladder and pilocarpine for the tracheal and urinary bladder smooth muscles.

Oxotremorine had very low activity on the uterus. The stereoselectivity of muscarinic ACh receptors (mAChRs) for cis-L(+)-and cis-D(-) dioxolane was low in the urinary bladder and uterus and high in the ileum and trachea. Most antagonists showed little selectivity between different organs, but S(-)-phenylcyclohexylglycoloyl choline was 6 times more active on the urinary bladder than on the ileum and AF-DX 16 was 12 30 times more active on the atria than on the smooth muscles. Among the N alkyl derivatives of benzilylcholine, the octyl derivative as 400 times more active on the ileum than on the atria, while among the N alkyl derivatives of ONB, the N-decyl derivative was 41 times more active on the ileum than on the potency of various agonists and their stereolsomers on differences in the potency of various agonists and their stereolsomers on different smooth muscles cannot be explained by differences in the accessibility of receptors or in receptor reserve. (ABSTRACT TRUNCATED AT 250 MORDS)

21 MEDLINE on STN 93103664 MEDLINE PubMed ID: 1281645 L12 ANSWER 53 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER:

PubMed ID: 1281045 Pogsible regulatory role of dynorphin A in the wrinary bladder. Berggren A; Dahlstrom A; Rubenson A; Sillen U Department of Pediatric Surgery, Ostra Hospital, AUTHOR -

CORPORATE SOURCE: Journal of neural transmission. General section, (1992) 90

SOURCE: (1) 33-44. Journal code: 9002201. ISSN: 0300-9564.

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: Austria Journal; Article; (JOURNAL ARTICLE)

English Priority Journals FILE SEGMENT:

ENTRY MONTH:

ENTRY DATE:

GUAGE: English
E SEGMENT: Priority Journals
RY MONTH: 199301
RY DATE: Entered STN: 19930122
Last Updated on STN: 19950129
Entered Medline: 19930127
Muscle Strips from rat and human detrusor were studied using indirect immunofluorescence and electrical nerve stimulation in an organ bath.
Immunoreactivity towards dynorphin was observed in varicose nerve fibres in the detrusor muscle and around immunonegative nerve cell bodies in the prevesical ganglia of the rat. In vitro dynorphin A (1-13)
(10(-13)-10(-6) M) strongly facilitated detrusor contraction induced by electrical field stimulation (EFS). This facilitation was counteracted by morphine (10(-10) and 10(-8) M) and naloxone (10(-10) and 10(-8) M) in a competitive manner. The facilitation could also be counteracted by the addition of the kappa-receptor antagonist M(n) 2266 (10(-7) M).
Muscarinic blockade, achieved with atropine (10(-6) M), did not alter the effect of dynorphin A (1-13). Addition of phentolamine mesylate (10(-6) M), and propranolol (10(-6) M) per se facilitated the EFS-induced contractions. Both adrenergic blockade as well as the addition of the substance P blocker spantide, counteracted the facilitating effect of dynorphin A (1-13). In conclusion: Dynorphin A immunoreactive material was found to be present in nerves in the rat detrusor and in prevesical ganglia. Dynorphin A (1-13) facilitated the detrusor contraction, possibly via actions on kappa-opioid receptors and interaction with non-cholinergic nerves.

L12 ANSWER 55 OF 121 ACCESSION NUMBER: 9: MEDLINE on STN DOCUMENT NUMBER:

21 MEDLINE On STN
93059868 MEDLINE
PubMed ID: 1433649
Effect of tetrodotoxin on the phasic and tonic responses of
isolated rabbit urinary bladder smooth
muscle to field stimulation.
Tammela T I; Wein A J; Levin R M
Division of Urology, University of Pennsylvania School of
Medicine, Philadelphia.
P50-DK 39257 (NIDDK)
(NIDDK)

AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER:

RO-1-DK 26508 (NIDDK) RO-1-DK 33559 (NIDDK)

SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

Medicine, Philadelphia.

TRACT NUMBER: P50-DK 39257 (NIDDK)

R0-1-DK 26508 (NIDDK)

RCE: JOURNAL of urology, (1992 Dec) 148 (6) 1937-40.

JOURNAL COMENTY: United States

JOURNAL ARTICLE)

GUAGE: English

E SEGMENT: Abridged Index Medicus Journals; Priority Journals

RY MONTH: 199212

RY DATE: Entered STN. 19930122

Entered Medicine; 19921233

The response of the rabbit wrinary bladder to field

stimulation (80 volts, 2-12 Hz, 1 meer duration) is biphasic, consisting of an initial phasic contraction mediated by cholinergic and purinergic neurotramentters, followed by a prolonged tonic contraction which is solely cholinergic. Obstructive hypertrophy of the bladdar to natively of contraction in the conic component of the contractive response as compared to the phasic component. This results in a severe dysfunction in the ability of the bladdar to maintain tension and empty efficiently may be related to a degeneration of nerves innervating the bladdar smooth muscle. In addition to the well documented neuropathy, the bladdar undergoes hypertrophy /- hyperplasia of both smooth muscle and interstitial cellular elements, alterations in the metabolism of substrates, alterations in the appearance of structural and smooth muscle proteins, and alterations in the deposition of collagen. The purpose of this study was to 1) to create a specific neuropathy in the absence of the additional structural, smooth muscle, and metabolic changes that are induced by partial outlet obstruction; and 2) determine if the contractile dysfunctions induced by the neuropathy was induced in leal assume that are induced by partial outlet obstruction. In the present study, a progressive "smooth muscle, and metabolic changes that are induced by partial outlet obstruction and 2) determine if the contractile dysfunctions induced by the neuropathy was induced in leal assume that provide a strips of male rabbit wrinary bladder smooth muscle by incubating isolated strips of unlared by a percent and the present study, a progressive "smooth muscle neuropa

ANSWER 55 OF 121 MEDLINE on STN (Continued) significantly greater than the ED50 following ATP desensitization. This may indicate that there are separate synaptic elements for cholinergic and purinergic transmission. (ABSTRACT TRUNCATED AT 400 WORDS) L12 ANSWER 55 OF 121

L12 ANSWER 57 OF 121 ACCESSION NUMBER: 9 MEDLINE on STN 92303044 MEDLINE PubMed ID: 1819171 DOCUMENT NUMBER: PubMed ID: 1819171
[Acetylcholinesterase and the ADH-dependent transport of water in the amphibian bladder].
Atsetilkholinesteraze i ADG zavisimyi transport vody v mochevom puzyre amfibii.
Bagrov Ia Iuu Manusova N B; Ostretsova I B
Tsitologiia; (1991) 33 (11) 141 52.
Journal code: 0417363. ISSN: 0041-3771.
USSR TITLE: AUTHOR: SOURCE: PUB. COUNTRY: DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE) Russian Priority Journals 199207 LANGUAGE: FILE SEGMENT:

ENTRY DATE:

NGUAGE: Russian

THEY MONTH: 199207

Last Updated on STN: 19920731

Last Updated on STN: 19920731

Last Updated on STN: 19920731

It was found that acetylcholine (ACh) at the concentration of 10(-3) M inhibited ADH-stimulated water transport through the wall of amphibian urinary bladder. This effect was suggested to be caused by an interaction of ACh was completely suppressed in the presence of various AChE inhibitors (physostigmine, proserine, armine, 6d-42, acridine iodmethylate), while an inhibitor of butyrylcholinesterase (BChE), AD-4, failed to affect it. In accord with this observation the activity of AChE (but not of BUChE) was demonstrated in the urinary bladder epithelium. Since, in addition to the hydrosmotic effects of pituitrine, 8-arginine-vasopressin or oxytocin, ACh blocked also effects of forskolin or cyclic AMP, one may conclude that it acts at some post-cyclic AMP production stage. AChE-dependent inhibition of the ADH-stimulated water transport decreased significantly when the serosal pH was raising from 7.2 to 8.0, but was augmented by serosal acidification (pH 6.8), whereas such pH alterations did not affect the activity of the epithelium AChE. The effect of ACh under consideration was suppressed by adding amiloride (10(4) M) to the serosal solution. Similarly, the ACh effect was blocked by an inhibitor of Ca-dependent H-channels, 4-aminopyrdine, which in addition prevented the inhibition of the ADH-stimulated water transport by the serosal acidification. It was noteworthy that some other K+ channel blockers (Ba2+, Cs+, tetraethylammonium, apamine, quinine) did not affect either the water transport or the antipituitrine effect of ACh interaction with AChE. It is unlikely that the acidification is merely a consequence of the ACh hydrolysis, rather the ACh-AChE interaction induces directly an increase in the proton conductivity of the basolateral membrane of the urinary bladder epithelium.

L12 ANSWER 56 OF 121 MEDLINE on STN ACCESSION NUMBER: 92395838 MEDLINE PubMed ID: 1381764 DOCUMENT NUMBER:

TITLE:

CORPORATE SOURCE:

PubMed ID: 1381764
Selectivity of class I antiarrhythmic agents, disopyramide, pirmenol, and pentisomide for peripheral muscarinic
M2 and M3 receptors.
Endou M, Hattori Y: Gando S; Kanno M
Department of Pharmacology, Hokkaido University School of Medicine, Sapporo, Japan.
Journal of cardiovascular pharmacology, (1992 May) 19 (5) 674-81.
Journal code: 7902492. ISSN: 0160-2446.
United States SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

Outrial; Article; (JOURNAL ARTICLE) English Priority Journals 199210

LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

ENGRANT: English
ESGGMENT: Priority Journals
RY MONTH: 199210
RY MONTH: 199210
Last Updated on STN: 19921023
The interactions of the class I antiarrhythmic agents, disopyramide, pirmenol, and pentisomide with peripheral muscarinic receptors were investigated by binding assay with [3H]N-methylscopolamine ([3H]NMS) as a ligand. All the agents inhibited the specific [3H]NMS binding to membrane preparations obtained from quinea pig submandibular gland (SG) and urinary bladder (UB) smooth muscle.
The competition curves of these agents for [3H]NMS binding to SG membranes were monophasic, indicating competition with [3H]NMS at a single site. Comparison of results with those of our previous binding experiments using quinea pig left atrial (LA) membranes, showed that pirmenol had sevenfold lower affinity for glandular-type muscarinic receptors (M3) than for cardiac-type muscarinic receptors (M3). On the other hand, the dissociation constants (Ki) for disopyramide and pentisomide in SG were comparable to the high affinity Ki values for these agents at M2 receptors. The competition curves of the three agents for [3H]NMS binding to UB membranes were biphasic and showed high- and low-affinity states of binding. The high- and low-affinity K1 values for pirmenol in UB were similar to its K1 values at M2 and M3 receptors obtained in LA and SG, respectively. The high affinity K1 values for dispyramide and pentisomide were consistent with the respective K1 values determined in SG, whereas the low-affinity binding sites for these agents were presumably the result of their allosteric interactions with the receptors. All agents at higher concentrations slowed the dissociation of (3H]NMS elicited by an excess of atropine in both UB and SG, thus indicating allosteric interactions. (ABSTRACT TRUNCATED AT 250 MORDS)

L12 ANSWER 58 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR .

All MEDLINE On STN
92299927 MEDLINE
PubMed ID: 1607601
Evidence for inhibitory nicotinic and
facilitatory muscarinic receptors in cholinergic
nerve terminals of the rat wrinary
bladder
bladder
Somogyi G T; de Groat W C
Department of Pharmacology, University of Fittsburgh,
Pennsylvania 13261.
DK 37241 (NIDDK) CORPORATE SOURCE:

CONTRACT NUMBER: D DK 42369 (NIDDK) MH 30915 (NIMH)

SOURCE: Journal of the autonomic nervous system, (1992 Feb) 37 (2)

Journal code: 8003419. ISSN: 0165-1838.

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

RCE: Journal of the autonomic nervous system, (1992 Feb) 37 (2) 89-97.
Journal code: 8003419. ISSN: 0165-1838.

COUNTRY: Netherlands
UMENT TYPE: Journal, Article; (JOURNAL ARTICLE)
UMCAGE: English
E SEGMENT: Priority Journals
RY MONTH: 199207
RY DATE: Entered STN: 19920731
Last Updated on STN: 19920731
Entered Medline: 19920722
Cholinergic prejunctional modulatory receptors on parasympathetic nerves in the rat urinary bladder were studied by measuring
3H-acetylcholine (Ach) release in muscle strips from the bladder body. Electrical field stimulation markedly increased 3H-Ach overflow in strips preloaded with 3H-choline. Oxotremorine (1 microM), an MZ receptor agonist and DMPP (10 microM) a nicotinic (N) receptor agonist indicating the presence of three types of modulatory receptors. The anticholinesterase agent, physostigmine in concentrations of 1, 5 and 25 microM and meostigmine (5 microM) increased Ach release (44-7104). However a low concentration of physostigmine (0.05 microM) decreased release. Pirenzepine, an ML muscatinic antagonist or atropine blocked the increased Ach release in physostigmine-treated strips, but in normal strips pirenzepine did not change release and atropine increased release. (3764 and 3914 respectively) in physostigmine-treated strips. The response to McN-3 343 was blocked by pirenzepine -Arbbocurarine (DTC), a nicotinic receptor blocker, enhanced Ach release in the presence of physostigmine but proved to be ineffective in normal preparations. Thee findings suggest that all three cholinergic receptors (MI facilitatory, N inhibitory) and MZ inhibitory are activated by endogenous Ach in physostigmine treated preparations. The will be important in future studies to determine whether MI and MZ mechanisms can also be activated under more physiological conditions in the bladder and whether they are present at other cholinergic

L12 ANSWER 59 OF 121
ACCESSION NUMBER: 91
DOCUMENT NUMBER: POTITLE: (

21 MEDLINE on STN
92183654 MEDLINE
PUMMed ID: 1724648
(+/-)-Terodiline: an MI-selective muscarinic
receptor antagonist. In vivo effects at muscarinic
receptors mediating urinary bladder
contraction, mydriasis and salivary secretion.
Noronha-Blob L; Prosser J C; Sturm B L; Lowe V C; Enna S J
NOVA Pharmaceutical Corporation, Baltimore, MD 21224.
European journal of pharmacology, (1991 Aug 29) 201 (2.3) AUTHOR: CORPORATE SOURCE: SOURCE:

135-42. Journal code: 1254354. ISSN: 0014-2999.

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

Netherlands Journal; Article; (JOURNAL ARTICLE)

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

English Priority Journals

SUAGE: English E SEGMENT: Priority Journals PT MONTH: 199204 E SEGMENT: Priority Journals PT MONTH: Entered STN: 19920426 E SEGMENT: Priority And selectivity of racemic terodiline (N text.butyl-1-methyl-3,3-diphenylpropylamine HCl) for muscarinic receptor subtypes was determined from functional responses of rabbit vas deferens (M1), guinea pig atria (M2) and bladder detrusor muscle (M3). Guinea pig atria (M2) and bladder detrusor muscle (M3). Selective for Mi relative to M2 (11-fold) and M3 (19-fold) receptors. Like pirenzepine, (*/-)-terodiline does not distinguish between M2 and M3 receptors in vitro. The peripheral actions of (*/-) terodiline were evaluated in vivo in terms of its ability to induce mydriasis, and to inhibit salivary secretion and urinary bladder contraction. (*/-)-Terodiline given s.c. was equipotent in inhibiting intravesical bladder pressure and carbachol-induced salivary secretion (1D50 = 59 mg/kg). These results suggest that the in vivo actions of racemic terodiline at (M3) receptors mediating bladder contraction may not be separable from its actions at receptors mediating mydriasis and salivation. Moreover, its effects on the pupil and salivary glands are apparently not mediated through M1 receptors. Together, these findings help clarify the action of (+/-)-terodiline in the treatment of neurogenic bladder

L12 ANSWER 61 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: OF 121 MEDLINE ON STN ER: 92070820 MEDLINE R: PubMed ID: 1959843

Pubmed ID: 199843 Pharmacological analysis of drug interactions of disopyramide and its congeners with peripheral muscarinic acetylcholine receptors.

AUTHOR: CORPORATE SOURCE:

Endou M
Department of Pharmacology, Hokkaido University School of

SOURCE:

Medicine, Sapporo, Japan. [Hokkaido igaku zasshi] Hokkaido journal of medical science, (1991 Sep) 66 (5) 677-93. Journal code: 17410290R. ISSN: 0367-6102.

Japan

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: Journal; Article; (JOURNAL ARTICLE) Japanese Priority Journals

FILE SEGMENT:

ENTRY MONTH ENTRY DATE:

Japanese E SEGMENT: Priority Journals RY MONTH: 199201

Externed Medline: 1992010

The interactions of the antiarrhythmic agents, disopyramide (D) and its congeners, pirmenol (Pr) and pentisomide (Pt), with peripheral muscarinic receptors (m-AchR) were investigated using binding and functional assays. D, Pr and Pt inhibited the specific binding of [3H]-N-methyl scopolamine ([3H]-NMS) to membrane fractions prepared from guinea pig left atria (LA), submandibular glands (SG) and vrinsry bladdars (UB) in a concentration-dependent warner. Computer-assisted analysis showed that the displacement curves with D obtained from LA and UB were shallow and beat fitted by a two-site model, whereas D interacted with a single class of binding sites in SG. Kinetic experiments measuring [3H]-MNS dissociation revealed the existence of allosteric interaction of D with m-AchR, and it might be responsible for the low affinity components of the displacement curves in LA and UB. The pKi values for D in high-affinity receptor sites in LA and UB (pKH) were very close to the pKi for D obtained in SG, and corresponded well to the pAz values of around 6.0 for antagonism against the carbachol-induced mechanical responses of LA and UB. Pt interacted with m-AChR with qualitatively very similar fashion to that of D, but its potency was very weak (1/10 of D). Pr interacted with a single class of binding sites in LA and SG with pKi of 6.02 and 5.18, respectively, indicating that the affinity of Pr to glandular m-AChR (M3) was 7 fold lower than that to cardiac one (M2). The displacement curve with Pr in UB was best fitted by a two-site model with pKi of 5.93 and pKL of 5.20. The pAZ for Pr in LA and UB were 6.47 and 5.55, respectively, suggesting the existence of a mixed population of M2 and M3 in UB and the contribution of M3 to its concluded that Pr is able to distinguish M2 from M3 and that D and Pt have almost similar affinity to both subtypes of m-AChR. Pr was less potent than D in interaction with M3.

L12 ANSWER 60 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 92131842 MEDLINE
DOCUMENT NUMBER: 1775512
Effects of pregnancy on muscarinic receptor
density and function in the rabbit wrinary
bladder.

Dladder. Levin R M; Zderic S A; Ewalt D H; Duckett J W; Wein A J Division of Urology, Hospital of the University of Pennsylvania, Philadelphia. RO-1 DR26508 (NIDDK) AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER:

RO-1 DK33559 (NIDDK)

RO1 DK39086 (NIDDK)

SOURCE: Pharmacology, (1991) 43 (2) 69-77. Journal code: 0152016. ISSN: 0031-7012.

PUB. COUNTRY: Switzerland

Journal; Article; (JOURNAL ARTICLE) English DOCUMENT TYPE:

LANGUAGE:

FILE SEGMENT: Priority Journals ENTRY MONTH:

ENTRY DATE:

Description of the response of the rabbit urinary bladder to component of the response of the shader and purinergic response of the physiology and pharmacology of the urinary bladder to field stimulation is cholinergic and 40% is purinergic. Although the purinergic response represents a significant proportion of the initial (phasic) pressure response to field stimulation is cholinergic and 40% is purinergic. Although the purinergic response represents a significant proportion of the initial (phasic) pressure response to field stimulation of the isolated whole bladder, it contributes only 10 15% of the ability of field stimulation to empty the bladder. The current study investigates the effects of pregnancy on the contractile responses of the isolated urinary bladder to cholinergic and purinergic stimulation. The results of these studies indicate that pregnancy induces substantial changes in the physiology and pharmacology of the urinary bladder. The following data are consistent with the theory that pregnancy substantially increases the relative purinergic component of the response to field stimulation (and presumably neuronal stimulation); (1) there was a significantly greater response of the bladders isolated from pregnant rabbits to low-frequency field stimulation; (2) atropine was more effective at inhibiting the pressure generation of bladders isolated from virgin female rabbits; (4) the response of the bladders from pregnant rabbits to bethanechol was significantly reduced, whereas the response to ATP was significantly increased. In addition to these effects of pregnancy on bladder physiology, pregnancy induced a 50% decrease in the mucarinic receptor density of the urinary bladder body, which correlated very well with the 50% decrease in the contractile response to bethanechol.

MEDLINE on STN 91339907 MEDLINE PubMed ID: 1651865 L12 ANSWER 62 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

PubMed ID: 1651865
Effect of extracellular Ca2+ on cholinergic, KCl and phorbol ester-mediated phosphoinositide turnover and guinea pig urinary bladder contraction.
Lowe V C, Noronha-Blob L
Nova Pharmaceutical Corporation, Baltimore, MD 21224.
European journal of pharmacology, (1991 Mar 26) 195 (2) 273-9. TITLE .

AUTHOR: CORPORATE SOURCE:

SOURCE:

273-9. Journal code: 1254354. ISSN: 0014-2999. Netherlands Journal; Article; (JOURNAL ARTICLE) English Priority Journals PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
SY MONTH: 199107
Last Updated on STN: 19911013
Entered Medline: 19910920
The effect of extracellular Ca2+ ([Ca2+]o) on cholinergic, KCl and phorbol ester-mediated detrusor contractions was related to phosphoinositide (PI) breakdown in guinea pig urinary bladder. Carbachol (1.0 mM) elicited a 20 fold increase in inositol phosphate (IP) accumulation both in presence and absence of [Ca2+]o yielding the same EC50 value (approximately 12 microM). In contrast, carbachol-induced detrusor contractions were reduced by 35% without [Ca2+]o, but maximal efficacy was restored with Ca2+ replenishment. In absence of (Ca2+]o, repeated cholinergic stimulation yielded contractions only if tissues were intermittently equilibrated in (Ca2+]o. High K+ and PDBu evoked [Ca2+]o-dependent contractions. Ca2+ channel antagonists and divalent metal cations inhibited high K+ more potently than carbachol-mediated contractions. Together, these findings suggest multiple sources of Ca2+ for urinary bladder contraction, where voltage-sensitive responses depend primarily on (Ca2+)o and PI-linked mucarinic responses involved Ca2+ mobilization from intracellular stores as well. Clinical agents used for the treatment of urinary incontinence inhibited both carbachol-induced PI turnover and muscle contraction with the same rank order of potency both in presence and absence of [Ca2+]o. These findings suggest that the cholinergic mechanism of action of these agents involves the PI-Ca2+ effector system.

L12 ANSWER 63 OF 121
ACCESSION NUMBER: 91
DOCUMENT NUMBER: PUTITLE: CC

AUTHOR

21 MEDLINE on STN
91279839 MEDLINE
Pubmed ID: 2057522
Comparison of the contractile and metabolic effects of
muscarinic stimulation with those of KCl.
Levin R M; Hypolite J; Longhurst P A; Wein A J
Division of Urology, University of Pennsylvania School of
Medicine, Philadelphia.
P-50-DK 39257 (NIDDK) CORPORATE SOURCE:

CONTRACT NUMBER:

RO-1-DK 26508 (NIDDK) RO-1-DK 33559 (NIDDK)

NHIDMAN
Pharmacology, (1991) 42 (3) 142-50.
Journal code: 0152016. ISSN: 0031-7012.
Switzerland SOURCE:

PUB. COUNTRY

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT:

English Priority Journals

ENTRY MONTH:

ENTRY DATE:

SEGMENT: Priority Journals
(X MONTH: 199108)

Last Updated on STM: 19910818

Last Updated on STM: 19910818

Last Updated on STM: 19910801

Urinary bladder emptying is mediated primarily by a coordinated contraction of the bladder body in response to parasympathetic stimulation and muscatinic receptor activation.

In previous studies we presented evidence that the contractile response to bethanechol stimulation could be dissociated from the metabolic response through the use of diltiazem (calcium channel blockade). The conclusion from these studies was that muscatinic stimulation resulted in a significant increase in metabolic activity which was not directly associated with contraction. KCl stimulates contraction in isolated strips by directly depolarizing the membrane rather than from binding to specific membrane receptors. The current study directly compares the metabolic and contractile activity of bethanechol (muscatinic stimulation) with KCl (direct membrane depolarization). Isolated strips of rabbit urinary bladdar body were monitored in vitro for changes in intracellular-free calcium, NABH/NAD ratio, and contraction. Intracellular-free calcium was monitored by preincubation of isolated bladdar smooth muscle strips with FURA-2 AM and continuously measuring the fluorescence with an MBz surface spectrofluorometer using excitation wavelengths of 340 and 380 mm, and an emission wavelength of 510 mm. The NABH/NAD ratio was monitored with the MB2 surface spectrophotometer using an excitation wavelength of 366 mm and an emission wavelength of 450 mm. The NABH/NAD ratio was monitored with the MB2 surface spectrophotometer using an excitation wavelength of 366 mm and an emission wavelength of 450 mm. The NABH/NAD ratio arapid increase in intracellular-free calcium, and a slower increase in contractile force transducer connected to a Grass model D polygraph. The results can be summarized as follows. (1) Both bethanechol and KCl stimulated as sharp decrease in the NABH/NAD ratio, a rapid increase in intracellular-fre

L12 ANSWER 65 OF 121 MEDLINE on STN
ACCESSION NUMBER: 91192889 MEDLINE
DOCUMENT NUMBER: Pubmed ID: 1672862
TITLE: DuF 753 is a specific antagonist for the angiotensin

receptor. Rhaleb N E; Rouissi N; Nantel F; D'Orleans-Juste F; Regoli

Department of Pharmacology, Medical School University of CORPORATE SOURCE:

Sherbrooke, Quebec, Canada. Hypertension, (1991 Apr) 17 (4) 480-4. Journal code: 7906255. ISSN: 0194-911X. United States PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE: English Priority Journals

SEMENT: Priority Journals
IY MONTH: 199105
IX DATE: Entered STN: 19910602
Last Updated on STN: 19980206
Entered Medline: 19910513
2-n-Butyl-4-chloro-5-hydroxy-methyl-1-(2'-(1H)-tetrazol-5-yl)biph enyl-4-yl)methyllimidazol potassium salt (DuP 753) is a nompeptide angiotensin II receptor antagonist that inhibits the contractile effects of angiotensin II competitively and shows pA2 values of 8.27 on the rabbit aorta and jugular vein, 8.66 on the rat portal vein and stomach, 8.19 on the rat wrinary bladder. This agent (more than 10(-5) M) exhibits no agonistic activity and does not affect the contractile effects of norepinephrine, acetylcholine, bradykinin, desArg9-bradykinin, substance P, neurokinin A, neurokinin B, or bombesin in the various tissues. The present results demonstrate that DuP 753 is a pocent nompeptide antagonist with high affinity, specificity, and selectivity for the angiotensin receptor.

L12 ANSWER 64 OF 121 ACCESSION NUMBER: 9

21 MEDLINE on STN
91259787 MEDLINE
PubMed ID: 1828510
Biochemical and functional characteristics of
bladder muscarinic receptors and effects
of experimental diabetes in rats.
Morita T DOCUMENT NUMBER: TITLE:

Morita T Department of Urology, Akita University School of Medicine. Nippon Hinyokika Gakkai zasuhi. Japanese journal of urology, (1991 Jan) 82 (1) 52-60. Journal code: 2984841R. ISSN: 0021-5287. CORPORATE SOURCE: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: Japan Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: FILE SEGMENT: Japanese Priority Journals

ENTRY MONTH: ENTRY DATE:

SURGE: Japanese
E SECHENT: Priority Journals
RY MONTH: 199107
RY DATE: Entered STN: 19910802
Entered STN: 19910802
Entered Month: 19910716

Bladder dysfunction is a common complication of diabetes
mellitus and is attributed in part to peripheral neuropathy. Voiding
function is mainly controlled by musearinic receptor function.
Therefore, I investigated first the biochemical and functional
characteristics of urinary bladder musearinic
receptors and then the effects of experimental diabetes on them.
Experimental diabetes was induced in 2 month-old male rate by intravenous
injection of 65 mg/kg of streptozotocin (STZ). Effects of diabetes
mellitus were investigated 2, 4 and 8 weeks after injection of STZ. The
amount of musearinic receptors labelled with 3H-quinclidinyl
bensylate (QNB) was higher in the bladder dome of diabetic
animals than control animals, while the affinity for its binding sites was
similar in both groups. Musearinic agonists and antagonists
inhibited 3H-QNB binding with similar inhibitory
constants (Ki) in control and diabetic domes. The rank order of
inhibition of 3H-QNB binding by musearinic agonists and
antagonists: bethanechol greater than pirenzepine greater than
carbamylcholine greater than acetylcholine greater than atropine, is
consistent with the absence of Mi receptors in the bladder dome.

In functional studies musearinic agonists induced a larger
contractile response in bladder dome muscle strips from 8
week-old diabetic animal than those from receptor binding studies.
These data show a direct correlation between the diabetes—induced
biochemical and functional alterations in musearinic receptor
properties of the rat bladder.

ANSWER 66 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

MEDLINE on STN
91132462 MEDLINE
PubMed JD: 1993995
Enantioners of oxybutynin: in vitro pharmacological characterization at M1, M2 and M3 muscarinic receptors and in vivo effects on urinary bladder contraction, mydriasis and salivary secretion in guines pigs.
Noronha-Blob L; Kachur J F
Nova Pharmaceutical Corporation, Baltimore, Maryland.
Journal of pharmacology and experimental therapeutics, (1991 Feb) 256 (2) 562-7.
Journal code: 0376362. JSSN: 0022-3565.
United States
Journal; Article; (JOURNAL ARTICLE)

AUTHOR:

CORPORATE SOURCE:

PUB. COUNTRY: Onited Scates Journal; Article; (JOURNAL ARTICLE) English Priority Journals 199103

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

Segment: Priority Journals
Y MONTH: 199103
Entered STN: 19910405
Last Updated on STN: 19970203
Entered Medline: 19910321
The major side effects of racemic oxybutynin (OXY), which is used in the treatment of urinary incontinence are dry mouth (xerostomia) and blurred vision (mydriasis). Righly purified enantioners of OXY [(R)OXY, (S)OXY] were compared with the racemate both in vitro in functional studies and in vivo in guinea pigs to evaluate their pharmacological action relative to their adverse effects. The affinity of (R)OXY and (S)OXY for different muscarinic receptor subtypes was determined using field stimulated rabbit vas deferens (M1) and guinea pig atria (M2) or bladder (M3) strips. Stereoselective antimuscarinic effects [(R)OXY greater than or equal to (R/S)OXY wore slightly more evident at all three receptor subtypes; the isomeric ratio [(S)OXY/(R)OXY] were evident at all three receptor subtypes; the isomeric ratio [(S)OXY/(R)OXY] exception (2 4 fold, P less than .01) for M1 and M3 relative to M2 muscarinic receptors. Stereoselectivity was also evident in vivo for volume-induced urinary bladder contractions as measured by cystometrogram parameters [(S)OXY/(R)OXY approximately 11, mydriasis [(S)OXY/(R)OXY approximately 136] and salivary gland secretory responses [(S)OXY/(R)OXY approximately 136] and salivary gland secretory responses [(S)OXY/(R)OXY approximately 136] and salivary gland secretory in the minimum second contractions of (R)OXY and (R/S)OXY for mydriasis and salivation were similar to those for inhibition of intravesical bladder pressure. Also, (R)OXY and (R/S)OXY dequipotently antagonized cholinergic-mediated CNS effects in mines. Collectively, the data suggest that the activity of (R/S)OXY raceded predominantly in the (R)-enantioner. However, it appears that (R)OXY and (R)OXY and first principal therapeutic and side effect profile.

L12 ANSWER 67 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: TITLE

21 MEDLINE on STN
91081462 MEDLINE
PubMed ID: 2175437
Muscarinic cholinergic antibody in experimental
autoimmune myocarditis regulates cardiac function.
Paaerez Leiros C; Sterin-Borda L; Cossio P; Borda E S
Centro de Estudios Farmacologicos y Botanicos (CEFYBO)
(ex CEFAPRIN), CONICET, Buenos Aires, Augentina.
Proceedings of the Society for Experimental Biology and
Medicine. Society for Experimental Biology and Medicine
(New York, N. Y.), (1990 Dec) 195 (3) 356-63.
Journal code: 7505892. ISSN: 0037-9727. CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY United States

Journal; Article; (JOURNAL ARTICLE) English DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: Priority Journals ENTRY MONTH:

ENTRY DATE:

SEGMENT: Priority Journals
Y MONTH: 199101
Y DATE: Entered STN: 19910322
Last Updated on STN: 19910322
Entered Moltime: 19910129
Evidence is presented showing that in experimental autoimmune myocarditis, there are certain components in IgG fraction of the sera that bind to myocardium muscarinic cholinergic receptors. The autoimmune IgG simulated the biologic effect of cholinergic agonists because (i) it increased CGMP levels, (ii) it decreased CAMP stimulated levels, and (iii) it reduced heart contractility and diminished reactivity to exogenous acetylchoine. Autoimmune IgG inhibited the binding of specific muscarinic cholinergic radioligand to purified myocardial membranes behaving as noncompetitive inhibitors. The recognition appears to be organ specific because the autoimmune IgG did not bind to muscarinic cholinergic receptors of urinary bladder. The presence of antibody in living myocardial cells might be related to some of the immunopathologic mechanisms participating in the pathogenesis of the experimental autoimmune myocarditis.

ANSWER 69 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

21 MEDLINE on STN
91067580 MEDLINE
PubMed ID: 2251221
Characterization of muscarinic cholinergic
receptors in membrane preparations from rat prostatic
-dancervicoms

adenocarcinoma.

Batra S; Christensson P I; Hartley-Asp B
Department of Cancer Pharmacology, Pharmacia LEO
Therapeutics AB, Helsingborg, Sweden.
Prostate, (1990) 17 (4) 261-8.

Journal code: 8101368. ISSN: 0270-4137.
United States
Journal, Article; (JOURNAL ARTICLE) CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

English

LANGUAGE: FILE SEGMENT: Priority Journals ENTRY MONTH: ENTRY DATE:

E-SEGMENT: Priority Journals
YY MONTH: 199101
RY MONTH: 199101
RY DATE: Entered STN: 19910308
Last Updated on STN: 19970203
Entered Medline: 19910117
The binding characteristics of 3H quinuclidinyl benzilate (QNB) to muscarinic sites in isolated plasma membrane fractions from R-3327
Dunning tumors (H and AT-1 sublines); ventral, dorsolateral prostate; and urinary bladder of the rat were studied. QNB binding to all preparations, except from AT-1 tumors, was specific, saturable, and of high affinity. The AT-1 tumors completely lacked specific (QNB binding. The muscarinic receptor density in H tumors was twofold and twentyfold higher than that in the ventral prostate and dorsolateral prostate respectively. The receptor density in the urinary bladder benefit of the urinary bladder. QNB binding in H tumors was strongly inhibited by classical muscarinic receptor antagonists carbacholine and pilocarpine. In contrast to scopolamine or atropine, inhibition by pirenzepine and AF-DXII6 was relatively low. These data indicate that the muscarinic receptor in Dunning H tumors is of M3 type.

ANSWER 68 OF 121 MEDLINE on STN 91077650 ACCESSION NUMBER: DOCUMENT NUMBER:

POTOTOS MEDLINE PUDMed ID: 2257437 High and low-affinity binding sites for [3H] alpha, beta-methylene ATP in rat uxinary bladder

AUTHOR CORPORATE SOURCE:

SOURCE

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE

High and low-affinity binding sites for [3H] alpha, beta-methylene ATP in rat urinary bladder membranes.

HOR: BO X N; BURNSTOCK G
DEPARTE SOURCE: Department of Anatomy and Developmental Biology, University College London.

RCE: British journal of pharmacology, (1990 Oct) 101 (2) 291-6.
JOURNAY: ENGLAND: United Kingdom UNENT TYPE: JOURNAL ARTICLE)

BUAGE: ENGLAND: United Kingdom ENGLAND: University Orders Type: JOURNAL ARTICLE)

BUAGE: ENGLAND: United Kingdom English ENGLAND: University Journals

RY MONTH: 199101

RY DATE: Entered STN: 19910322
Last Updated on STN: 19970203
Entered Medline: 199103101

1. The characteristics of (3H)-alpha, beta-methylene adenosine
5'triphosphate ((3H)-alpha, beta MeATP) binding to membrane preparations of rat urinary bladder detrusor were studied. 2. The rat bladder membrane preparation was obtained by multiple centrifugation. (3H)-quinuclidiny benzilate ((3H)-(NB)) binding to this preparation demonstrated that the muscarinic receptor density was 4.32 times higher than that in the homogenate. (3H) alpha, beta-MeATP binding was increased 3.88 times. 3. Saturation analysis revealed that the rat bladder membrane contained a high density of (3H)-alpha, beta MeATP binding sites, which could be divided into a high affinity component (Kd = 8.1-8.9 nM) and a low-affinity binding was reduced from 10.32 pmol mg·l protein in magnesium-free buffer to 4.62 pmol mg·l protein with 25 mM MgCl2, while the maximum binding in a concentration-dependent manner. The maximum binding in a reduced from 58.84 pmol mg·l protein to 14.24 pmol mg·l protein. Kd values were not greatly affected. 5. The binding was a rapid reversible process. The association rate constants were 7.64 x 10(7) M·l min-1 for high-affinity binding, and 7.31 x 10(6) M·l min-1 for low-affinity binding, and 7.31 x 10(6) M·l min-1 for low-affinity binding, and 7.31 x 10(6) M·l min-1 for low-affinity binding, and 7.31 x 10(6) M·l min-1 for low-affinity binding, and 7.31 x 10(6) M·l min-1 for low-affinity binding, and 7.31 x 10(6) M·l mi

L12 ANSWER 70 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: P

TITLE:

AUTHOR : CORPORATE SOURCE.

SOURCE

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE

ANSWER 70 OF 121 MEDLINE on STN
ESSION NUMBER: 90384910 MEDLINE
DUMENT NUMBER: PubMed ID: 2402477
LB: Effects of some antidepressants on the volume-induced reflex contractions of the rat urinary bladder: lack of correlation with muscarinic receptors affinity.
HOR: Pleta C; Poggest E; Angelico P; Quarneri L; Testa R
Pharmacological research official journal of the Italian Pharmacological research official journal of the Italian Pharmacological research official journal of the Italian Pharmacological Society, (1990 Jul Aug) 22 (4) 421-32.
JOURNAL COUNTRY: BNOLAND: United Kingdom
JUNGEN: PROPER SOURCE: PROPER SOURCE: PROPER SOURCE: Provincy Journals SEGMENT: Priority Journals
TY DATE: Entered STN: 19901122
Last Updated on STN: 1990203
It has been suggested that tricyclic antidepressants such as imipramine, might exert their anti-enureric action by a blockade of muscarinic receptors in the detrusor muscle of the urinary bladder.
We have therefore investigated the effects of two tricyclic (imipramine and nortriptyline) and three atypical (citalopram, amineptine and mianserin) antidepressants on the micturition reflex and muscarinic receptors in rats. The micturition reflex and muscarinic receptors in rats. The micturition reflex pathway was monitored indirectly by recording the rhythmic intravesical pressure waves which occurred when the bladder was distended and maintained under constant saline-volume. The activity of the antidepressants was correlated to their potencies as antagonists of (3H)QNB binding to rat brain (mainly MI receptors) and bladder (mainly MI receptors) membranes, as well as antagonists of carbachol-induced contractions of rat bladder strips. Only imigramine and citalopram dose dependently inhibited the voiding contractions, whereas nortriptyline, imipramine and misnaerin (in order of potency) were active both in binding the micturition reflex. The present data seem to suggest that affinities for muscarinic receptors are unrelated to the inhibition of micturition reflex.

L12 ANSWER 71 OF 121 ACCESSION NUMBER: 9 DOCUMENT NUMBER: PO 1 MEDLINE on STN 90315725 MEDLINE PubMed ID: 2369797

runged 10: 2369797 Interaction of antiestrogens with binding sites for muscarinic cholinergic drugs and calcium channel

blockers in cell membranes Batra S

AUTHOR: CORPORATE SOURCE:

Department of Obstetrics and Gynecology, University

SOURCE

Department of Obstetrics and Gymecology, University Hospital, Lund, Sweden, Cancer chemotherapy and pharmacology, (1990) 26 (4) 310-2. Journal code: 786519, ISSN: 0344-5704. GERMANY, WEST: Germany, Federal Republic of Journal, Article, (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: English

Priority Journals FILE SEGMENT: 199008

ENTRY MONTH: ENTRY DATE:

SECONETT: Priority Journals
YM MONTH: 19900 STN: 19900 921
Last Updated on STN: 19900 921
Last Updated on STN: 19900 921
The interaction of tamoxifen and clomifene with membrane binding sites for the cholinergic ligand quinuclidinyl benzilate (QNB) and the dihydropyridine calcium antagonist nitrendipine was studied. Both tamoxifen and clomifene competed with [3H]-QNB and [3H]-nitrendipine for their binding to the receptor in the membrane fractions from the urlnary bladder and myometrium. The extent of inhibition as judged by the Ki values for both antiestrogens was similar at both receptor sites. The data suggest that the antiproliferative effects of tamoxifen may involve not only the intracellular estrogens receptor system but also receptors for neurotransmitters and membrane calcium channels.

L12 ANSWER 73 OF 121 ACCESSION NUMBER: 90 DOCUMENT NUMBER: PO MEDLINE on STN 90250642 MEDLINE PubMed ID: 2338651 PubMed ID: 2338651
Autonomic receptors in urinary tract: sex and age differences.
Latifpour J; Kondo S; O'Hollaren B; Morita T; Weiss R M Section of Urology, Yale University School of Medicine, New Haven, Connecticut. DK 38311 (NIDDK)
Journal of pharmacology and experimental therapeutics, (1990 May) 253 (2) 661-7.
Journal code: 0376362. ISSN: 0022 3565.
United States AUTHOR: CORPORATE SOURCE: CONTRACT NUMBER: PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: United States Journal; Article; (JOURNAL ARTICLE) English Priority Journals ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
SY MONTH: 199006
YE DATE: Entered STN: 19900720
Last Updated on STN: 19900720
Last Updated on STN: 19900720

Entered Medline: 19900621
As age and sex affect the function of the lower urinary tract, we studied the characteristics of adrenergic and cholinergic receptors in various parts of lower urinary tract smooth muscle of young (6 months) and old (4 1/2-5 years) male and female rabbits. Saturation experiments performed with [JH]prazosin, [JH]ychimbine, [3H]dihydroalprenolol and [JH]quinuclidinyl bensylate in rabbit bladdar base, bladdar dome and urethra indicate the presence of regional, sex- and age-related differences in the density of alpha-1, alpha-2, and beta adrenergic and muscarinic cholinergic receptors. Alpha 2 adrenergic receptor density is considerably higher in the female than in the male urethra of both age groups, whereas the higher density of beta adrenergic receptors in the female than in the male bladdar base is observed only in the younger animals. The density of muscarinic receptors is higher in bladdar dome than in bladdar base or urethra in young rabbits of both sexes. In the old animals, the density of muscarinic receptors in the ladder base increases to the level observed in bladdar dome hand the density of muscarinic receptors in bladdar adrenergic receptors in the urethra and beta adrenergic receptors in the bladdar dome and bladdar the urethra and beta adrenergic receptors in the bladdar dome and bladdar the urethra and beta adrenergic receptors in the bladdar dome and bladdar base and similar in both sexes and at both ages. Beta-2 a drenergic receptors are shown to be predominant in bladdar base and bladdar dome of rabbits. Parallel studies in rabbit urethra, adult rat cortex and neonatal rat lung show that the urethra slipha 2 adrenergic receptors are of the alpha-2A subtype.

ANSWER 72 OF 121 MEDLINE on STN 90277703 MEDLINE PubMed ID: 2161851 ACCESSION NUMBER: DOCUMENT NUMBER:

Atropine resistant transmission in partially depolarized rat urinary bladder. Carpenter F TITLE:

AUTHOR CORPORATE SOURCE:

Carpenter F. Department of Pharmacology, University of Alabama, Birmingham 35294.
Journal of autonomic pharmacology, (1990 Apr) 10 (2)

SOURCE: 97 107.

Journal code: 8106455, ISSN: 0144-1795, PUB. COUNTRY: ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

DOCUMENT TYPE: LANGUAGE:

English Priority Journals FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

SIGNAGE:

SIGNAMENT:

Priority Journals

Y MONTH:

199007

Last Updated on STN: 19900824

Entered Medline: 19900716

1. Phasic contractile responses of the intact rat urinary

bladder to the muscarinke agonists carbachol and
pilocarpine became nearly blocked as the concentrations were progressively
increased to 200 500 microm. In contrast, tonic contractile responses

remained elevated throughout progressive increases in agonist
concentration. 2. Nerve-induced phasic contractions to 1 Hz stimuli were
potentiated throughout progressive increases in the concentration of
muscarinic agonists. However, these responses were more atropine
sensitive than untreated controls and responses to 1 Hz stimuli were
nearly abolished. 3. After inhibition of cholinestrame
, the action of cholinergic transmitter released during prolonged nerve
stimulation may extend to the tonic contractile state of the
bladder and potentiate responses to 1H stimuli. Nerve-induced
responses were more atropine sensitive than untreated controls. 4.
Bladder tone was increased and nerve-induced contractions to 1 Hz
stimuli were also potentiated in an elevated K. environment. However,
atropine sensitivity of nerve-induced responses w s reduced. 5.

Nerve-induced bladder contractions were linked to the tonic
contractile state of the bladder muscle, controlled
physiologically by muscarinic receptors. Since phasic
contractile responses to muscarinic agonists were abolished at
high concentrations by receptor desensitization, nerve-induced responses
must be elicited under these conditions by a non-cholinergic transmitter.

L12 ANSWER 74 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR: CORPORATE SOURCE:

SOURCE

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

ANSWER 74 OF 121 MEDLINE on STN

ESSION NUMBER: 90165615 MEDLINE

JUMENT NUMBER: 90165615 MEDLINE

LE: Mechanism of action of nicotine in isolated urinary

bladder of guinea-pig: involvement of tachykinin(s)

released by nicotine in the drug's sympathomimetic effect.

HOR: HISAYAMA T; Shinkai M; Takayanagi I; Toyoda T

PORATE SOURCE: Popartment of Chemical Plantmacology, Toho University School

of Pharmaceutical Sciences, Chiba, Japan.

RCE: Archives internationales de pharmacodynamie et de therapie,

(1989 Sep Oct) 301 277-84.

JOURNAL ARTICLE)

GUAGE: English

ES EEMBERT: Priority Journals

RY MONTH: 199003

RY DATE: Entered STN: 1990019

A sympathetic neurone blocking drug, guanethidine, and a tachykinin

antagonist, [D-Argl, D-Pro2, D-Trp7,9, Leull]-substance P (rpwwl-SP),

partially inhibited the contractile response to nicotine to the

same degree in the isolated detrusor strips of guinea pig urinary

bladder. Application of rpwwl-SP completely abolished the

inhibitory effect of guanethidine on the nicotine-induced

contraction, suggesting that the tachykinin(s)-ergic transmission might be

involved in the sympathemientic effect of nicotine. Conversely, when the

preparation was created with guanethidine to block release of a mediator

from the sympathetic nerve, the inhibitory effect of ripwdl-SP

was diminished, suggesting an exclusive contribution of the sympathetic

nerve communications to the action of the tachykinin(s)

suggested that nicotine may release acetylcholine and AP to contract the

detrusor strips, and that acetylicholine output may be increased by an

unknown substance released from the sympathetic nerve by nicotine. In

preparations treated with atropine, rpwwl-SP had no effect on the

nicotine-induced contraction. The concentraction-response curves for

carbacinal and AP twee noi influenced by rpwwl-SP. After tachyphylaxis to

capsaicin developed, the nicotine-induced contraction was not affected.

It is suggested that in guinea-pig detrusor, tachykinin(s) from

coppalating and the fa

L12 ANSWER 75 OF 121 ACCESSION NUMBER: 90 MEDLINE on STN DOCUMENT NUMBER:

AEDLINE ON STN
90130540 MEDLINE
PubMed ID: 2613733
Effects of selective cholinergic antagonists and
alpha, beta-methylene ATP on guinea-pig urinary
bladder contractions in vivo following pelvic nerve

AUTHOR: CORPORATE SOURCE:

Dladde Contractors stimulation. Peterson J S; Noronha-Blob L Nova Pharmaceutical Corporation, Baltimore, Maryland

SOURCE:

Nova Pharmaceutical Corporation, Baitimore, Maryland 21224 2788 Journal of autonomic pharmacology, (1989 Oct) 9 (5) 303-13. Journal code: 8106455. ISSN: 0144-1795. BNGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY:

DOCUMENT TYPE:

English Priority Journals

LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

SEMMENT: Priority Journals
Y MONTH: 199003
Y DATE: Entered STN: 19900328
Last Updated on STN: 19900328
Last Updated on STN: 19900305
1. An in vivo preparation measuring functional detrusor muscle strength in terms of intravesical bladder pressure (Pves) following in situ pelvic nerve stimulation has been developed in urethane-anaesthetized guinea pigs. 2. The increase in bladder pressure following pelvic nerve stimulation was abolished by topical lidocaine or tetrodotoxin, suggesting a neurogenic origin for the in vivo contractile response. 3. Cholinergic antagonists (i.v.) decreased the amplitude of the peak pressure response by about 508 at both high (30 Hz) and low (5 Hz) stimulation rates, with a rank order of potency of atropine greater than propantheline greater than nexhipdrosiladifenidol greater than nivenzepine greater than hexahydrosiladifenidol greater than nivenzepine greater than methoctramine. 4. The P2 purine receptor antagonist, alpha, beta methylene ATP (i.v.), antagonized pelvic nerve stimulated bladder contractions differentially at 5 and 30 Hz. At low frequencies, alpha, beta-methylene ATP was both more potent (2.5-fold) and more efficacious (-77 compared to -55% delta) than at 30 Hz. Atropine and alpha, beta-methylene ATP together completely inhibited the contractile response. 5. Together, the findings indicate that in guinea pigs, urinary bladder contractions induced by pelvic nerve stimulation in vivo may be mediated by both muscarinic and purinergic receptors and that these bladder contractions may be mediated by the MZ beta subtype rather than by M1 or M2 alpha

ANSWER 77 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR:

CORPORATE SOURCE:

21 MEDLINE on STN

90074839 MEDLINE
PubMed ID: 2480167
Multiple sources of calcium for contraction of the human
urinary bladder muscle.
Maggi C A; Giuliani S; Patacchini R; Turini D; Barbanti G;
Giachetti A; Meli A
Pharmacology Department, Res. Labs., A. Menarini
Pharmaceuticals, Plorence, Italy.
British journal of pharmacology, (1989 Nov) 98 (3) 1021-31.
Journal code: 7502536. ISSN: 0007-1188.
ENGLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

COMMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

KGLAGE: English

E SEMANT: Priority Journals

ENG MONTH: 199001

ENGY DATE: Entered STN: 1990028

Last Updated on STN: 19900208

Last Updated on STN: 19900208

1. KCl, carbachol, neurokinin A and endothelin produced concentration dependent contractions of mucosa-free muscle strips from the dome of the human uxinary bladder. The maximal response to carbachol or neurokinin A exceeded that to KCl, while the maximal response to endothelin approached that to KCl. 2. Nifedipine (1 microM) abolished the response to KCl but had little or no effect on the response to carbachol or neurokinin A exceeded that to KCl. 2. Nifedipine (1 microM) markedly potentiated the response to KCl but had little or no effect on the response to KCl but had little or no effect on the response produced by the other stimulants. 3. Superfusion of the strips with a nominally calcium (Ca)-free medium containing EDTA (1 mM) for 30 min markedly preduced the response to carbachol, neurokinin A and endothelin, although a small response was still evident at high concentrations. Likewise, after a prolonged (60 min) superfusion of the strips with a high K (80 mM) Ca-free medium plus EDTA (1 mM) these three agonists still produced a small contractile response. 4. The nifedipine (1 microM) resistant response to carbachol, neurokinin A or endothelin was markedly depressed by Lacl3 (1 mM). In contrast, the nifedipine-(1 microM) resistant response to carbachol was not modified by Nicl2 (0.1 mM) or omega-conotoxin (0.1 microM). 5. Caffeine produced divergent effects depending upon the temperature of incubation: a relaxation at 37 degrees C. The latter was markedly inhibited by procaine (3 mM) but unaffected by nifedipine (1 microM). 6. After a prolonged (60 min) superfusion with a high K, Ca-free medium containing EDTA the response to carbachol (100 microM) was abolished by previous exposure to procaine (3 mM). Conversely, the response to endothelin (1 microM) was unaffected by procaine. The response to endo

21 MEDLINE ON STN 90095937 MEDLINE PubMed ID: 2600844 Enkephalinergic inhibition in parasympathetic ganglia of the urinary bladder of the cat. ANSWER 76 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

AUTHOR:

cat.

de Groat W C; Kawatani M

Department of Pharmacology, School of Medicine, University

of Pittsburgh, PA 15261.

AM 316888 (NIADDK) CORPORATE SOURCE:

CONTRACT NUMBER: A MH 30915 (NIMH) NS 25254 (NINDS)

Journal of physiology, (1989 Jun) 413 13-29. Journal code: 0266262. ISSN: 0022-3751. ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY

DOCUMENT TYPE: LANGUAGE:

English Priority Journals FILE SEGMENT:

ENTRY MONTH:

SUAGE: English
E SSCMENT: Priority Journals
EN MONTH: 199001
Entered Medline: 19900128
Last Updated on STN: 19970203
Entered Medline: 19900129
1. Repetitive stimulation (10-20 Hz, 0.5-5 s duration) of the preganglionic nerves to ganglia on the surface of the urinary bladder of the cat produced a prolonged inhibition
(duration, 30-65 s) of the postganglionic action potentials, elicited by low-frequency stimulation (0.25-1 Hz) of another preganglionic nerve to the same ganglion. 2. Intra-arterial administration of naloxone, an opiate antagonist (20-50 micrograms/kg), reduced the magnitude and duration of this heterosynaptic inhibition and also blocked the depression of ganglionic transmission elicited by the intra-arterial administration of leucine-enkephalin (0.1 10 micrograms/kg). 3. Naloxone did not alter adrenergic inhibition elicited by repetitive stimulation of the hypogastric nerve or exogenous noradrenaline. Naloxone did not alter the postganglionic firing elicited by single stimuli or trains of low-frequency (1 3 Hz) stimuli to the preganglionic nerves. 4. Heterosynaptic inhibition was not altered by the administration of antagonists for alpha-adrenergic (dihydroergotamine, praxosin, yohimbine), muscarinic (atropine), purinergic (theophylline) or GABAcrgic (picrotoxin) receptors. 5. A delta-selective opiate receptor agonist, DSLET (D-Ser2-leucine-enkephalin Thr), inhibited parasympathetic ganglionic transmission in low doses (mean threshold dose, 0.02 microgram/kg, 1.A.), whereas a mu-opiate receptor agonist, morphine sulphate, produced only a small depression in larger doses (mean threshold dose, 0.02 microgram/kg, 1.A.). Bthylketocyclaxocine, which has an affinity for kappa-receptors did not alter transmission in relatively large doses (1 mg/kg, 1.A.). 6. These findings coupled with previous immunocytochemical demonstrations of leucine-enkephalin like immunocytochemical demonstrations of leucine-enkephalin like immunocytochemical demonstrations of leucine-enkephalin like immunocytochemical demonstratio

L12 ANSWER 78 OF 121 ACCESSION NUMBER: 90 DOCUMENT NUMBER: PO

MEDLINE on STN
90064117 McDLINE
PubMed ID: 2573724
Reorganization of sympathetic preganglionic connections in
cat bladder gasglia following parasympathetic
denervation.
de Groat VC (Kawatani M
Department of Pharmacology, University of Pittsburgh, PA
15261.
AM 316888 (NIADDK)

ROHTLIA

CORPORATE SOURCE:

CONTRACT NUMBER: A MH 30915 (NIMH) NS 25254 (NINDS) SOURCE: J

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

TRACT NUMBER: AM 316888 (NIADDK)
MH 30915 (NIMH)
NS 25254 (NINDS)
NS 25254 (NINDS)
RCE: Journal code: 0266262. ISSN: 0022-3751.
. Journal code: 0266262. ISSN: 0022-3751.
. COUNTRY: ENGLAND: United Kingdom
UMENT TYPE: Journal; Article; (JOURNAL ARTICLE)
GUAGE: English
E SEGMENT: Priority Journals
RY MONTH: 199001
RY DATE: Entered STN: 19900328
Last Updated on STN: 19970203
Entered Medline: 19900105
1. Experiments were undertaken to examine the mechanisms involved in the reorganization of sympathetic efferent pathways to the urinary bladder of the cat following chronic unilateral, parasympathetic preganglionic demervation or on the normally innervated side of the bladder in unilaterally demervated preparations elicited low-amplitude (10-25 cmHzO) transient (10-30 s) bladder contractions and non synaptic axonal volleys on bladder postganglionic nerves. However, after chronic (3-22 months) sacral preganglionic demervation, the propagation enerve stimulation on the side of the demervation elicited large (60-80 cmH2O) and more sustained (4 5 min) bladder contractions as well as synaptically mediated firing on bladder postganglionic nerves. 3. The vesicoexcitatory effects of hypogastric nerve stimulation on the chronically denervated side were not altered selectively by the adrenergic blocking agent, phenoxybenzamine, but were blocked by atropine and hexamethonium suggesting that the responses were mediated by muscarinic and nicotinic cholinergic synapses. These drugs did not influence the responses elicited by hypogastric nerve stimulation on the horonically denervated side were not altered selectively by the adrenergic blocking agent, phenoxybenzamine, but were blocked by atropine and hexamethonium suggesting that the responses were mediated by muscarinic and nicotinic cholinergic synapses. These drugs did not influence the responses elicited by hypogastric nerve stimulation of the pelvic and hypogastric nerves on one side of the bladder. 4. Following more extensive chronic unilateral intext pelvic nerve elicited po

L12 ANSWER 79 OF 121
ACCESSION NUMBER: 9
DOCUMENT NUMBER: P

21 MEDLINE on STN
90017077 MEDLINE
PubMed ID: 2477832
Contractile properties of human prostate adenomas and the development of infravesical obstruction.
Gup D I; Shapiro E; Baumann M; Lepor H
Division of Urologic Surgery, Jewish Hospital of St. Louis,

AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER:

PUB. COUNTRY: DOCUMENT TYPE:

DIVISION OF Urologic Surgery, Jewish Ho MO 63110. RR05491-25 (NCRR) Prostate, (1989) 15 (2) 105-14. Journal code: 8101368. ISSN: 0270-4137. United States Journal: Article; (JOURNAL ARTICLE)

English

LANGUAGE: FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE:

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

ANSWER 81 OF 121 MEDLINE ON STN
SSION NUMBER: 89279738 MEDLINE
MENT NUMBER: 90400 ID: 2542813

**Muscarinic receptors: relationships among phosphoinositide breakdown, adenylate cyclase inhibition, in vitro detrusor muscle contractions and in vivo cystometrogram studies in guinea pig bladder.

**North Robert Strands Blob L. Lowe V. Batton M. Committee B. Committee B.

bladdar.
Noronha Blob L; Lowe V; Fatton A; Canning B; Costello D; Kinnier W J
Nova Pharmaceutical Corporation, Baltimore, Maryland.
Journal of pharmacology and experimental therapeutics, (1989 Jun) 249 (3) 843-51.
Journal code: 0376362, ISSN: 0022-3565.

CORPORATE SOURCE: SOURCE:

United States Journal; Article; (JOURNAL ARTICLE)

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY DATE:

MENT TYPE: Journal; Article; (JOURNAL ARTICLE)

JUNGE: English

S EMEMENT: Priority Journals

YY MONTH: 198907

Last Updated on STN: 19980206

Entered Medline: 19890721

The relationships between activation of muscarinic receptors in guines pig bladder measured as carbachol stimulated inositol phosphate (IP) accumulation, oxotremorthe-induced adenylate cyclase (AC) inhibition and bladdar detrusor smooth muscle contraction determined in vitro as well as in vivo in the slow filling cystometrogram (CMG), were analyzed from the potencies of a number of muscarinic antagonists to block these responses. Significant positive linear correlations were found among the inhibitory potencies of 10 muscarinic antagonists to inhibit phosphoinositide (PI) turnover and both detrusor muscle contraction in vitro, as well as peak intravesical bladder pressure in vivo in the CMG (r = 0.8, P less than .01). In contrast, there was no significant correlation between the potency of antagonists to block the AC inhibitory response and either in vitro or in vivo guinea pig bladder contractions (P greater than .05). Muscarinic agonists inhibited basal AC activity to a maximum of 20% in a GTP-dependent, Na+-sensitive manner and dose dependently stimulated both PI breakdowm (3- to 4 fold) and isolated detrusor contractions. Again, a significant correlation (r = 0.9, P less than .01) was calculated among the potencies of seven muscarinic agonists to elicit PI turnover and in vitro muscle contraction, whereas no significant correlation and presumably IP-induced Ca++ release may function as the transducing mechanism for cholinergic contraction of the urinary bladder. (ABSTRACT TRUNCATED AT 250 WORDS)

L12 ANSWER 80 OF 121 ACCESSION NUMBER: 96 MEDLINE on STN 90015245 MEDLINE PubMed ID: 2797217 DOCUMENT NUMBER:

TITLE:

CORPORATE SOURCE:

PubMed ID: 2797217
Prejunctional effects of muscarinic agonists on 3H-acetylcholine release in the rat urinary bladder strip.

D'Agostino G; Chiari M C; Grana E Institute of Pharmacology, Pavia, Italy.
Naunyn-Schmiedeberg's archives of pharmacology, (1989 Jul) 340 (1) 76-81.

Journal code: 0326264. ISSN: 0028-1298.
GERMANY, WEST: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals
198911
Entered STN: 19900228 SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE:

FILE SEGMENT: ENTRY MONTH

ENTRY DATE:

L12 ANSWER 82 OF 121 ACCESSION NUMBER:

DOCUMENT NUMBER:

21 MEDLINE on STN
89199462 MEDLINE
PubMed ID: 2539454
Contractile responses in bladder body,
bladder neck and prostate from rat, guinea pig and

cat. Cohen M L; Drey K Lilly Research Laboratories, Eli Lilly and Company, Lilly Corporate Center, Indianapolis, Indiana. Journal of pharmacology and experimental therapeutics, (1989 Mar) 248 (3) 1063-8. Journal code: 0376362. ISSN: 0022-3565. United States AUTHOR: CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE) English

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE:

SURGE: English

S. SEGMENT: Priority Journals

YMONTH: 198905

Last Updated on STM: 19900306

Entered Medline: 19890512

Lower wrinary tract smooth muscle displays marked heterogeneity
in pharmacologic responsiveness to contractile agents. The present study
details differences among species with regard to muscarinic,
adrenergic, histaminergic and serotonergic agonists in the bladdar
body, bladdar neck and prostate from guines pig, rat and cat.
Under in vitro conditions, all smooth muscle preparations contracted to
potassium chloride. The muscarinic agonist, carbamylcholine,
produced maximal contraction, whereas alpha receptor agonists exerted only
minimal, if any, effect in bladdar body preparations from all
three species. In contrast, alpha receptor mediated responses
predominated relative to muscarinic responses in bladdar
neck preparations from all three species. Prostatic contractility was
examined in tissue from guinea pig and rat and contraction occurred to
both alpha and muscarinic receptor agonists. Contractile
response to norepinephrine in bladdar neck and prostate was
potentiated by neuronal uptake inhibition but not by beta
receptor blockade. Serotonin and histamine exhibited more diverse effects
among species and tissues. In general, histamine contracted all three
tissues from guinea pig with minimal contraction occurring in tissues from
rat or cat. On the other hand, serotonin markedly contracted the cat
bladdar body and rat prostate, but exerted no effect on tissues from
the quinea pig. These data retinforce and detail the heterogeneity of
pharmacologic contractile responses in lower urinary tract
smooth muscle. Furthernore, the studies document the relative similarity
among species in cholmergic and adrenergic menonsiveness and the
dissimilarity among species in serotonergic and histaminergic

L12 ANSWER 83 OF 121
ACCESSION NUMBER: 85
DOCUMENT NUMBER: POTITLE: Me MEDLINE on STN

AUTHOR: CORPORATE SOURCE:

21 MEDLINE On STN
89150848 MEDLINE
PubMed ID: 3228673
Mechanism of action of nicotine in isolated urinary
bladder of guinea-pig.
Hisayama T; Shinkai M; Takayanagi I; Toyoda T
Department of Chemical Pharmacology, Toho University School
of Pharmaceutical Sciences, Chiba, Japan.
British journal of pharmacology, (1988 Oct) 95 (2) 465 72.
Journal code: 7502536. ISSN: 0007-1188.
ROLAND: United Kingdom
Journal; Article; (JOURNAL ARTICLE)
English

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE

Journal code: 7502536. ISSN: 0007-1188.

COUNTRY: ENOLAND: United Kingdom
JOURNALD: United Kingdom
JOURNALD: United Kingdom
JOURNALD: English
SEGMENT: Priority Journals
SEGMENT: Prior

ANSWER 85 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

21 MEDLINE on STN
89074918 MEDLINE
PubMed ID: 2904773
Regulation of [3H]GARA release from strips of guinea pig
urinary bladder.
Shirakawa J; Taniyama K; Iwai S; Tanaka C
Department of Aneethesiology, Kobe University, School of
Medicine, Japan.
American journal of physiology, (1988 Dec) 255 (6 Pt 2)
R888-93. TITLE:

AUTHOR: CORPORATE SOURCE:

SOURCE:

Journal code: 0370511. ISSN: 0002-9513. United States

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

onited States
Journal; Article; (JOURNAL ARTICLE)
English Priority Journals

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

E SEGMENT: Priority Journals
RY MONTH: 198901
RY DATE: Entered STN: 19900308
Last Updated on STN: 19950206
Entered Medline: 19890126
The presence of receptors that regulate the release of gamma-aminobutyric acid (GABA) was studied in strips of the guinea pig urinary bladder. GABA (10(-8)-10(-5) M) and muscimol (10(-8)-10(-5) M), but not baclofen (10(-5) M), reduced the Ca2+-dependent, tetrodotoxin-resistant release of 13H(GABA evoked by high K+ from the urinary bladder strips preloaded with [3H]GABA. The inhibitory effect of muscimol was antegonized by bicuculline and potentiated by diazepam, clonazepam, and pentobarbital sodium. The potentiating effect of clonazepam was antagonized by Ro 15-1788. Acetylcholine (Ach) inhibited the high K+-evoked release of [3H]GABA. The inhibitory effect of ACh was antagonized by atropine sulfate and pirenzepine but not by hexamethonium. Norepinephrine (NE) inhibited the evoked release of [3H]GABA. The inhibitory effect of NE was mimicked by clonidine, but not by phenylephrine, and was antagonized by yohimbine but not by prazosin. These results provide evidence that the release of GABA from strips of guinea pig urinary bladder is regulated via the blocuculline-sensitive GABAA receptor, MI-muscarinic, and alpha 2-adrenergic receptors.

ANSWER 84 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

21 MEDLINE On STN
85094334 MEDLINE
PubMed ID: 2453333
In vivo motor effects of substance P on the rat
urinary bladder.
Berggren A; Ahlman H; Dahlstrom A; Rubenson A; Sillen U
Department of Pediatric Surgery, East Hospital, Goteborg,
Sweden. ATITHOR CORPORATE SOURCE:

sweden.
Journal of neural transmission, (1988) 74 (3) 207-17.
Journal code: 0337042. ISSN: 0300-9564. SOURCE.

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: Austria Journal; Article; (JOURNAL ARTICLE)

English Priority Journals 198902 FILE SEGMENT: ENTRY MONTH:

SEGMENT: Priority Journals
YM MONTH: 198902
YM MONTH: 198902
Thered STN: 1990308
Thered STN: 1990308
Thered STN: 1990308
Thered Mediane: 19890219
Thered Mediane: 19890219
Thered Mediane: 19890219
Thered Mediane: 19890213
The motor control of this organ was evaluated. Regional injection of SP (0.4 modes i.a.) into the superior vesical artery elicited a prompt bladder contraction; this motor response was dosedependent. The detrusor contraction could be completely inhibited by a SP analogue, (D Pro2, D-Trp7,9)-SP (45-90 mmoles i.a.). Furthermore, the detrusor contraction evoked by preganglionic stimulation of the pelvic nerves was partially inhibited by the same antagonist in a higher dose (55% reduction at a total dose of 150-300 mmoles). The contractile response to SP (0.5 nmoles i.a.) was also significantly reduced after blockade of muscarinic receptors with atropine (50% reduction at 1 mg/kg i.a.) or after ganglionic blockade with hexamethonium (75% reduction at 25 mg/kg i.v. + 50 mg/kg hr i.a.). Immunocytochemical studies demonstrated the occurrence of SP-immunopositive nerve terminals in the detrusor part of the rat urinary bladder. Based on these findings it is suggested that SP may act as a neurotransmitter/modulator in this organ. The mechanism of action for SP on the detrusor seems to be complex and may involve ganglionic transmission via both types of cholinoceptors as well as direct activation of smooth muscle.

L12 ANSWER 86 OF 121 ACCESSION NUMBER: 8 DOCUMENT NUMBER: P

MEDLINE on STN
89014401 MEDLINE
PubMed ID: 3173349
Dissociation of the metabolic from the contractile response
to muscartnic stimulation in the rabbit
urinary bladder.
Ruggieri M R, Wein A J, Hypolite J A; Levin R M
Division of Urology, University of Pennsylvania School of
Medicine, Philadelphia.
RO-1-DK 33559 (NIDDK)
(NIDDK) TITLE

AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER: RO-1-DK-2-6508 SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT

DEVISION Of Urclogy, University of Pennsylvania School of Medicine, Philadelphia.

TRACT NUMBER: R0-1-DK 33559 (NIDDK)
RCE: Molecular and cellular biochemistry, (1988 Jun) 81 (2) 137-43.

Journal code: 0364456. ISSN: 0300-8177.

Netherlands
JOUREN: Netherlands
SEGMENT: JOURNAL ARTICLE)
JOURNAL ARTICLE)
JOURNAL BISH
SEGMENT: Priority Journals
EXT MONTH: 198811
The calcium dependence of contraction and NADH fluorescence was investigated in rabbit bladder stimulated with bethanechol or RCI. The absence of calcium in the bathing solution induced a rightward shift in the dose response to bethanechol for both contraction and NADH fluorescence. The contractic response was shifted to a greater degree than the fluorescence response and the maximal response to bethanechol was reduced by 80% for contraction but only 20% for NADH fluorescence. This rightward shift was also induced by the benzothiazepine calcium antagonist diltiazem (200 microM) and again the contractile response was shifted significantly more than the fluorescence response. The combination of zero calcium and 200 microM diltiazem virtually abolished contractions but only inhibited the NADH fluorescence by 65% at maximally effective bethanechol concentrations. Unlike the effect of diltiazem on the response to bethanechol, diltiazem calcom in the absence of any observable contractiie response. This metabolic response to be Manacrinic stimulation (decreased NADH) can occur in the absence of any observable contractile response. This metabolic response may be due to post receptor signal processing events. For KCI stimulation, the NADH response is probably secondary to and a result of the contractile response.

L12 ANSWER 87 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

MEDILINE ON SIN
88273235 MEDLINE
PubMed ID: 3392052
Differential effects of pertussis toxin on
muccatinic responses in isolated atria and smooth

UTTHOR

CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

MEMT NUMBER: PubMed ID: 3392052
E: Differential effects of pertussis toxin on muscarinic responses in isolated atria and smooth muscle.

GR: Eglen R M; Huff M M; Montgomery W W; Whiting R L; SCRATE SOURCE: Institute of pharmacology, Syntex Research, Palo Alto, California 94303
GCE: Journal of autonomic pharmacology, (1988 Mar) 8 (1) 29 37.

Journal code: 8166455. ISSN: 0144-1795.
ENSIAND: United Kingdom
JUAGE: SEMSENT: Priority Journals
19 MONTH: 198808
ENGIAND: United Kingdom
JUAGE: Entered STN: 19900308
Last Updated on STN: 20021218
Entered Medline: 19880817
1. The effect of pretreatment with pertussis toxin has been studied on responses to muscarinic agonists in guinea-pig atria and smooth muscle in vitro. 2, 48 h after a single intravenous injection of pertussis toxin (3.2-100 micrograms.kg-1), muscarinic receptor-mediated negative inotropic responses in the atria were inhibited in a dose-dependent manner, with complete abolition of responses occurring after administration of 100 micrograms kg-1. 3. In contrast, there was no effect on atrial positive inotropic responses to isoprenaline. In addition, no effect was observed on contractile responses to carbachol and pilocarpine in the ileum, trachea, oesophageal muscularis mucosea and urimary bladder, either in terms of potency or maximal response, at all dose levels of pertussis toxin studied. 4. It is concluded that muscarinic receptors in the atria, but not smooth muscle, are probably coupled to the inhibitory regulatory protein Ni, which is functionally inactivated by pertussis toxin. The differences in coupling between atrial and smooth muscle muscarinic receptors provide further evidence for muscarinic receptors in these two tissues.

L12 ANSWER 89 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

21 MEDLINE on STN
88242686 MEDLINE
PubMed ID: 3378565
Characterization of the muscarinic receptor
subtypes in the rat urinary bladder.
Monferini E; Giraldo E; Ladinsky H
Department of Biochemistry, Istituto De Angeli S.p.A., AUTHOR: CORPORATE SOURCE:

monnecini E; Giraldo E; Ladinsky H Department of Biochemistry, Istituto De Angeli S.p.A., Milan, Italy. Buropean journal of pharmacology, (1988 Mar 15) 147 (3) 453-8. SOURCE:

453-8. Journal code: 1254354. ISSN: 0014-2999. Netherlands

PUB. COUNTRY:

Netherlands Journal; Article; (JOURNAL ARTICLE) English Priority Journals 198807 DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

ENTRY MONTH:

ENTRY DATE:

E SEGMENT: Priority Journals
RY MONTH: 198807
RY DATE: Entered STN: 19900308
Last Updated on STN: 19900308
Entered Medline: 19880725
We investigated the nature of the muscarinic receptors present
in the rat urinary bladder by performing binding
studies with various selective (pirenzepine, AF-DX 116,
hexahydrostladitentidol, benshexol, 4-diphenyl-acetoxy N-methyl piperidine
methiodide, dicyclomine, secoverine) and classical (N-methylscopolamine,
atropine) antagonists. Competition experiments were carried out against
[JH]N-methyl scopolamine at 30 degrees C in Nai/Mg2+ HEPES buffer;
non-specific binding was determined in the presence of 1 microM
3-quinuclidinyl benzilate. Of all the antagonists examined, only AF-DX
116 exhibited a heterogeneous binding profile (nH less than 1).
Computer-assisted analysis showed that the data fitted best to a
two-binding site model, revealing the existence of high and low affinity
receptors. The affinity values of AF-DX 116, determined in binding
experiments carried out in heart and gland homogenates, allowed us to
classify the rat urinary bladder receptors into
cardiac and glandular subtypes. We suggest that the glandular receptor
subtype is involved in smooth muscle contraction, since AF-DX 116 was
equally potent in inhibiting smooth muscle contraction and the

L12 ANSWER 88 OF 121
ACCESSION NUMBER: 8
DOCUMENT NUMBER: PITITLE: PI

CORPORATE SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

PANSWER 88 OF 121 MEDLINE ON STN
PRESSION NUMBER: 88251531 MEDLINE
TUMENT NUMBER: PubMed ID: 2838033
TLE: Pharmacological activities of the main metabolite of flavoxate 3-methylflavone-8-carboxylic acid.

HOR: Cazulani P; Pietra C; Abbiati G A; Ceserani R; Oliva D; Civelli M; Tajana A; Nardi D

PORATE SOURCE: Research Division, Recordati S.p.a., Milan, Italy.
Arzneimittel-Forschung, (1988 Mar) 38 (3) 379-82.

Journal code: 0372660. ISSN: 0004-4172.

L. COUNTRY: Journal code: 0372660. ISSN: 0004-4172.

L. COUNTRY: Journal code: 0372660. ISSN: 0004-4172.

RESEMENT: Finity Journals
RY MONTH: 198807
RY DATE: Entered STN: 19900308
Entered Medine: 19880720
The pharmacological properties of 3-methylflavone-8-carboxylic acid (MPCA), the main metabolite of flavoxate, have been studied in vitro and in vivo. MFCA did not display antispasmodic activity on isolated organs contractions induced by histamine, acetylcholine or CaCl2, nor did it exhibit significant affinity for the rat brain alpha- and beta-adrenergic, serotonnic, muscarind, D2, opiate and Ca2+ receptors.

However, it showed a remarkable phosphodiesterase (PD) inhibiting activity. Moreover in vivo studies indicate an interesting activity of MFCA which inhibited the rat urinary bladder volume capacity and decreased micturition pressure in the rat cystometric recordings. The activity of MFCA win in the work in vivo experimental models, probably related to CAMP-FDE inhibitory properties, suggests that flavoxate's therapeutical potential might be partially sustained by its main metabolite.

L12 ANSWER 90 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 88172688 MEDLINE
DOCUMENT NUMBER: PUMMed ID: 2832620
Identification of receptor subtypes in the rabbit and human urinary bladder by selective radio-ligand

AUTHOR: CORPORATE SOURCE:

CONTRACT NUMBER: RO-1-DK-2-6508 SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

CLESION NUMBER: Pubmed ID: 232620

CLE: Winder Divided ID: 232620

CHOR: Levin R M, Ruggieri M R; Wein A J

Division of Urology, University of Pennsylvania School of Medicine, Philadelphia.

FRORT SOURCE: RO-1-DK-2-6508

CHORT RO-1-DK-2-6508

CHORT RO-1-DK-2-6508

CHORT SUMBER: RO-1-DK 33559 (NIDDK)

RCS: Journal of urology, (1988 Apr) 139 (4) 844-8.

Journal code: 0376374. ISSN: 0022 5347.

LOURDAY: United States

JOURNEY: United States

JOURNEY: Journal; Article; (JOURNAL ARTICLE)

CHORT: English

Abridged Index Medicus Journals; Priority Journals

ESEMENT: Abridged Index Medicus Journals; Priority Journals

RY DATE: English

ESEMENT: Abridged Index Medicus Journals; Priority Journals

RY DATE: Instead STN: 19900308

Recent advances in receptor technology have demonstrated that subtypes of each autonomic receptor exist. Using both direct radio-ligand studies and the inhibition of receptor binding by subtype selective pharmacological antagonists, we have studied the distribution of subtypes of alpha and beta adrenergic receptors and muscarinic cholinergic receptors in the urinary bladder of the rabbit and man. Alpha adrenergic receptors were quantified by direct binding of tritiated prazosin (alpha-1), yohimbine (alpha-2), and the non-selective alpha adrenergic receptors were quantified by the binhibition of the non-selective alpha adrenergic ligand dihydrocaproriptime (DHE). These studies demonstrated that the distribution of alpha receptor subtypes in the bladder base (for both rabbit and human) is approximately

80% alpha-1 and 20% alpha-2. Beta receptor subtypes were identified by the inhibition of The non-selective ligand 3H-dihydrocalprenalol

(DHA) by the beta-1 selective inhibitor ICI 89 and the beta-2 selective inhibitor ICI-118. Initial studies demonstrated hat the beta adrenergic density of the bladder body was 92 fmol per mg, protein for the rabbit and human. Inhibition of DHA binding by ICI-118 demonstrated as aligned class of receptor with an ICSO of approximately 9.0 microm for both rabbit and h

1.12 ANSWER 91 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

COMMENT:

ATTHOR

21 MEDLINE on STN
88034192 MEDLINE
PubMed ID: 3668160
Identification and characterization of muscarinic
cholinergic receptors in the human urinary
bladder and parotid gland.
Erratum in: J Auton Nerv Syst 1988 Mar;22(2):174. Bjorklund
A[corrected to Biorklund A]
Batra S; Biorklund A, Hedlund H; Andersson K E; Bjorklund A
AB Leo Research Laboratories, Helsingborg, Sweden.
Journal of the autonomic nervous system, (1987 Aug) 20 (2)
129-35. CORPORATE SOURCE: SOURCE:

Journal code: 8003419. ISSN: 0165-1838.

PUB. COUNTRY: Netherlands DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE) English

LANGUAGE: FILE SEGMENT:

Priority Journals

ENTRY MONTH: ENTRY DATE:

E SEGMENT: Priority Journals
NY MONTH: 198711

For DATE: Entered STN: 19900305

Last Updated on STN: 19900305

The binding characteristics of [3H]quinuclidinyl benzilate (QNB) to
muscarinic sites in isolated plasma membrane fractions of the
human urinary bladder and parotid gland were studied.
QNB binding to both preparations was of high affinity and low capacity.
Mean values for the apparent dissociation constants (Kd) for binding to
membrane preparations from the urinary bladder and
parotid glands were 22 and 34 pM and the Bmax values 234 and 456 fmol/mg
protein, respectively. Significance of difference between Kd and Bmax
values from the two tissues was at the level of P less than 0.005 and P
less than 0.05, respectively. QNB binding was inhibited by
muscarinic receptor antagonists with varying degree of
effectiveness. The mean values for the inhibition constant (Ki)
were significantly lower for oxybutynin, anirriptyline, and pirenzepine
but higher for secoverine in preparations of the urinary
bladder than of the parotid gland. The mean Ki values for
quinidine and verapamil were lower in the urinary
bladder than that in the parotid gland. Carbachol exhibited a
marked selectivity for the urlnary bladder (about
30-fold) compared with the parotid gland. The present data obtained in
two human tissues that are highly cholinergic in their innervation give
support to the argument for heterogeneity of the muscarinic

21 MEDLINE On STN 87304487 MEDLINE PubMed ID: 3622609 ANSWER 93 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

rounced ID: 3022007 Comparison of muscarinic acetylcholine binding in the urinary bladder and submandibular gland of the rabbit. Batra S TITLE:

AUTHOR:

European journal of pharmacology, (1987 Jun 12) 138 (1) SOURCE:

83-8. Journal code: 1254354. ISSN: 0014-2999. PUB. COUNTRY: Netherlands

Getnerianos Journal; Article; (JOURNAL ARTICLE) English Priority Journals 198710 DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

E SEGMENT: Priority Journals
RY MONTH: 198710

For DATE: Entered STN: 19900305

Last Updated on STN: 19970203

Entered Medline: 19871016

In order to explore the possibility of heterogeneity in peripheral muscarine receptors, receptors were characterized in membrane fractions isolated from rabbit urinary bladder and submandibular gland with [3H](DNB as radioligand, specific binding with very high affinity was found in both preparations. Although the BmaxS for binding in the two preparations were very similar, the mean KD value in the submandibular gland was significantly higher (P less than 0.005) than that in the bladder. Among the anticholinergic drugs, oxybutynin had a significantly lower value for the inhibition constant (Ki) in the submandibular gland whereas Ki for both secoverine and pirenzepine was significantly higher in this tissue than in urinary bladder. The Ki for carbacholine was about 7-fold higher in submandibular gland than in the bladder. Although quinidine and verapamil showed relatively weak binding to the muscarine receptor site, their Ki in the submandibular gland was significantly higher than that in the bladder. The results indicate that although there is a considerable similarity between muscarinic receptors in urinary bladder and submandibular gland, the differences in Ki values for different compounds in the two tissues support the argument favouring heterogeneity of muscarinic acetylcholine receptors in peripheral effector organs.

L12 ANSWER 92 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

AUTHOR SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

ANSWER 92 OF 121 MEDLINE ON STN
ESSION NUMBER: 87322496 MEDLINE
UNDERT NUMBER: PubMed ID: 3630733
LE: Source of calcium for contractions induced by
depolarization and muscarinic receptor
stimulation in rabbit urinary bladder.

HOR: Batra S; Siggern C; Anderson K E; Fovaeus M
RCE: Acta physiologica Scandinavica, (1987 Aug) 130 (4) 545-51.
Journal code: 0370352 ISSN: 0001-6772.
COLNTRY: ENGLAND: United Kingdom
UNDENT TYPE: Journal: Article; (JOURNAL ARTICLE)
GUACE: English
E SEGMENT: Priority Journals
RY MONTH: 198710
RY DATE: Entered Str.: 19900305
Entered Medline: 19871014
Omission of calcium or the inclusion of lanthanum in the bathing medium resulted in an almost complete inhibition of contractile
responses induced by either K* depolarization or carbachol in strips of rabbit urinary bladder. D-600 inhibited
K*-induced contractions significantly more than carbachol-induced responses. The influx of 45Ca into cells was stimulated both by K* depolarization and carbachol. Over a 2-min period the increase in 45Ca influx induced by high K* and carbachol was 98 and 65K, respectively.
Both lanthanum and D-600 blocked 45Ca influx stimulated by either K* depolarization or carbachol. The inhibition of carbachol during 45Ca efflux. These results indicate that the contractile responses of the urinary bladder to depolarization and to carbachol are selfux. These results indicate that the contractile responses of the urinary bladder to depolarization and to carbachol are shighly dependent on an extracellular source of calcium.

L12 ANSWER 94 OF 121 ACCESSION NUMBER: 8 DOCUMENT NUMBER: P

AUTHOR:

21 MEDLINE on STN
87198978 MEDLINE
PubMed ID: 3573166
Interaction between adrenergic and cholinergic nerve
terminals in the urinary bladder of
rabbit, cat and man.
Mattiasson A; Andersson K E; Elbadawi A; Morgan E; Sjogren

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

Mattiasson A; Andersson K E; Elbadawi A; Morgan E; Sjogren C
C
RCE: Journal of urology, (1987 May) 137 (5) 1017-9.
Journal code: 0376374. ISSN: 0022-5347.
COUNTRY: United States
JOURNAL ATTICLE)
JOURNAL ATTICLE)
JOURNAL ATTICLE)
JOURNAL ATTICLE)
MAGE: English
SEGMENT: Abridged Index Medicus Journals; Priority Journals
YF MONTH: 198706
MY DATE: Entered STN: 19900303
Last Updated on STN: 19900303
Entered Medline: 19870601
The influence of muscarfinic receptor stimulation (carbachol) and
blockade (scopolamine) on the release of 3H-labelled noradrenaline from
adhenergic neurons was investigated in isolated detrusor preparations from
rabbit, cat and man. A significant influence on the release of 3H from
adrenergic nerve terminals was found in the three species with a
concentration-dependent decrease and increase induced by carbachol and
scopolamine, respectively. Using the alpha 2-adrenoceptor stimulating and
blocking agents clonidine and rauvolscine in rabbit and human detrusor
preparations, the presence of prejunctionally located inhibitory
alpha 2-adrenoceptors could also be demonstrated. The findings indicate
the possibility of a functionally important interaction between
cholinergic and adrenergic nerves in the urinary bladder
mediated via inhibitory muscarinic receptors on
adrenergic nerve terminals.

L12 ANSWER 95 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

ALITHOR

21 MEDLINE on STN
87144723 MEDLINE
PUMMed ID: 2434871
Prostanoid synthesis by the rat urinary
bladdar: evidence for stimulation through muscarine
receptor-linked calcium channels.
Jeremy J Y, Mikhaildis D P, Dandona P
Naunyn-Schmiedeberg's archives of pharmacology, (1986 Dec)
344 (4) 463-7.
Journal code: 0326264. ISSN: 0028-1298.
GERMANY, MEST: Germany, Federal Republic of
Journal, Article: (JOURNAL ARTICLE)
English
Priority Journals
198704
Entered STN: 19900303

PUB COUNTRY

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

SEGMENT: Priority Journals

SEGMENT: Priority Journals

MONTH: 198704

Last Updated on STN: 19900303

Entered Medline: 19870420

An in vitro model for the study of muscarine receptor-mediated synthesis of prostacyclin (FGI2) and other prostanoids (FGE2 and FGF2 alpha) by the rat urinary bladder is described. FGI2 synthesis was stimulated by parasympathonimentic agents (carbachol greater than methacholine greater than arecoline; McNA 343, nicotine and dimethyl phenyl piperazinium were without effect). Methacholine (3 x 10(-6) mol x 1-1)-stimulated FGI2 synthesis was inhibited by muscarinic antagonists (atropine greater than ipratroprium bromide greater than gallamine greater than pirenzepine) and was completely abolished by the presence of ethylene diamine tetracetic acid (EDTA: 10 mmol X 1 1). Verapamil also inhibited methacholine stimulated FGI2 synthesis in a dose-dependent manner. The antagonistic action of atropine was shown to be competitive, but had no effect on calcium ionophore A23187-stimulated PGI2 synthesis. High concentrations of (k+) (up to 6.11 mol X 1-1) were without effect on FGI2 synthesis. FGE2, FGF2 alpha and FGI2 synthesis were all equally stimulated synthesis of these prostanoids was equally inhibited by atropine, ipratroprium bromide, gallamine, verapamil and EDTA. It is concluded that in vitro prostanoid synthesis by the rat urinary bladder: is stimulated by post ganglionic muscarine receptors; involves a muscarine receptor-linked calcium influx system; and is mediated by a predominance of M2 subtype receptors.

ANSWER 97 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

MEDLINE on STN
87036443 MEDLINE
PubMed 1D: 3772807
Presynaptic inhibitory muscarinic
receptors modulating [38] acetylcholine release in the rat
urinary bladder.
D'Agostino G; Kilbinger H; Chiari M C; Grana E
Journal of pharmacology and experimental therapeutics,
(1986 Nov) 239 (2) 522-8.
Journal code: 0376362. ISSN: 0022-3565.
United States
Journal; Article; (JOURNAL ARTICLE)
English
Priority Journals
198612
Entered STN: 1003

AUTHOR:

SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

E SEGMENT: Priority Journals
RY MONTH: 198612
RY DATE: Entered STN: 19900302
Last Updated on STN: 19900302
Entered Medline: 19961211
The occurrence of presynaptic muscarinic receptors
inhibiting the release of acetylcholine (Ach) from nerve terminals
was investigated in the rat urinary bladder. Strips
from the extractigonal area were preincubated with [3H]choline and
stimulated at 0.2 Hz. Both [3H]Ach and [3H]choline content were measured
in the rissue. The uptake of tritiated choline was prevented by
hemicholinium 3. The field stimulation at 2 Hz [360 shocks) produced a
release of tritium. Most of the induced outflow was found to be [3H]Ach.
Both tetrodotoxin treatment and calcium omission from the medium prevented
such an evoked outflow of tritium. When two electrical stimulations (S1
and S2) at 2 Hz [360 shocks) were carried out at 45-min intervals, an
S2/S1 ratio of 0.82 was found. Physostigmine reduced the evoked-release
of [3H]Ach whereas atropine increased it in a concentration-dependent
manner. Atropine antagonized the inhibitory effect of
physostigmine, so that the S2/S1 ratio did not vary significantly from
control experiments. Both carbachol and muscarine strongly decreased the
[3H]Ach evoked outflow. Muscarine increased the spontaneous outflow of
tritium also. These indings suggest that the urinary
bladder of the rat is equipped with presynaptic inhibitory
muscarinic receptors modulating ACh release from cholinergic
postganglionic neurons.

L12 ANSWER 96 OF 121 MEDLINE ON STN
ACCESSION NUMBER:
POCUMENT NUMBER:
BOCUMENT NUMBER:
CHOING PubMed ID: 3806496
Identification and characterization of muscarinic cholinergic receptors in the isolated plasma membranes and intact tissue of the urinary bladder.

PARTA S

**CONVERGENCE (1986) 6 (3-4) 227-46.

intact tissue of the urinary bladder.

Batra S
Journal of receptor research, (1986) 6 (3-4) 227-46.

Journal code: 8008358. ISSN: 0197-5110.

United States
Journal, Article; (JOURNAL ARTICLE)

English
Priority Journals

198702

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

Priority Journals
PRY MONTH: 198702
Entered STN: 19900303
Last Updated on STN: 19970203
Entered Medline: 19870224
The binding characteristics of 9H-quinuclidinyl benzilate (QNB) to strips of intact tissue and to isolated plasma membrane fraction (PM) from rabbit urinary bladder were studied. QNB binding to both preparations was of high affinity and low capacity. The equilibrium dissociation constants (KD) for binding to tissue strips and PM were 2.2 and 0.045 nM respectively. Muscarinic antagonists strips and PM were 2.2 and 0.045 nM respectively. Muscarinic antagonists
Ca-antagonist D-600, but not nifedipine caused an inhibition of QNB binding to PM. Vanadate, ouabain or N-ethylmelaimide had no significant effect on QNB binding. In contrast to the binding in PM, binding in the intact tissue was reduced by K-depolarization.

L12 ANSWER 98 OF 121 ACCESSION NUMBER: 80 DOCUMENT NUMBER: Pr

TITLE:

AUTHOR -

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

21 MEDLINE on STN
86282099 MEDLINE
PUMMed ID: 3755479
Action of pirenzepine on the human urinary
bladdar in vitro.
Zappia L; Cartella A; Potenzoni D; Bertaccini G
Journal of urology, (1986 Sep) 136 (3) 739-42.
Journal code: 0376374. ISSN: 0022 5347.
United States
Journal; Article; (JOURNAL ARTICLE)
English
Abridged Index Medicus Journals; Priority Journals
198609
Entered STN: 19900331 FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SAUMENT: Abridged Index Medicus Journals; Priority Journals; Y MONTH: 198609
Y DATE: Entered STN: 19900321
Last Updated on STN: 19800321
Entered Medline: 19860925
The novel compound pirenzepine was tested for its antimuscarinic effect on the human urinary bladder "in vitro." Its behavior towards the contractions induced by acetylcholine or bethanechol and towards electrically induced contractions was identical to that of atropine. However, its potency was 100 to 300 times lower than that of atropine. Results obtained with ganglion blocking agents, tetrodotoxin and cooled preparations of urinary bladder seem to indicate the virtually total absence of ganglionic cells. On the other hand they point out the fundamental role of post-synaptic muscarinic M2 receptors as the most important component of the cholinergic system in the bladder. Of course the existence of other transmitters released at the cholinergic nerve endings after electrical field stimulation cannot be excluded on the basis of our experiments.

L12 ANSWER 99 OF 121 ACCESSION NUMBER: 86 DOCUMENT NUMBER: P

AUTHOR:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE

ANSWER 99 OF 121 MEDLINE ON STN

SSTON NUMBER: 86034014 MEDLINE

MENT NUMBER: PubMed ID: 2414301

E: Urinary bladder by the use of agonists
and antagonists of the cholinoreceptors in the rat
urinary bladder by the use of agonists

AGR: Adami M; Bertaceini G; Coruzzi G; POli E

Journal of autonomic pharmacology, (1985 Sep) 5 (3)

197-205.

JOURNAI Of autonomic pharmacology, (1985 Sep) 5 (3)

197-205.

COUNTRY: ENOLAND: United Kingdom

MENT TYPE: Journal > Journal | Journal > Journal

L12 ANSWER 101 OF 121 ACCESSION NUMBER: 850 DOCUMENT NUMBER: Pub MEDLINE on STN

121 MEDLINS on STN
85058429 MEDLINE
PUMMed ID: 2981104
Effects of vasoactive intestinal polypeptide on isolated uterthral and wrinary bladder smooth muscle from rabbit and man.
Sjogren C: Andersson K E; Mattiasson A
Journal of urology, (1985 Jan) 133 (1) 136-40.
Journal code: 0376374. ISSN: 0022-5347.
United States
Journal; Article; (JOURNAL ARTICLE)
English
Abridged Index Medicus Journals; Priority Journals
198501
Entered STN: 1990320

AHTHOR

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SUAGE: English E SECMENT: Abridged Index Medicus Journals; Priority Journals BE SECMENT: Abridged Index Medicus Journals; Priority Journals BY MONTH: 198501

EN MONTH: 198501

EN FORTE: Entered STN: 19900320

Entered Medicus: 19850124

Vasoactive intestinal polypeptide concentration-dependently inhibited the contractant responses of isolated preparations of the female rabbit bladdar and urethra induced by electrical field stimulation and exogenous application of acetylcholine (bladdar) and noradrenaline (urethra). The inhibition of alpha-ademoceptor and muscarinic cholinoceptor-mediated activity in the urethra and bladdar amounted to 50 to 90 per cent of induced contractions. The nonadrenergic noncholinergic contraction induced by electrical field stimulation in the urethra was reduced slightly, whereas corresponding response in the bladdar was more sensitive. The maximum inhibition of both the electrically induced responses and contractions induced by exogenous noradrenaline and acetylcholine was of comparable size in the urethra and the bladdar. The effects of vasoactive intestinal polypeptide seemed to be exerted postjunctionally since no significant influence of the peptide was seen on the release of 31-horadrenaline from adrenergic nerve endings in the urethra. The effects of vasoactive intestinal polypeptide in human urethral and bladdar preparations were less consistent. The moradrenaline-induced contraction in urethral preparations was inhibited by 29-47-9 per cent (number 22). The effects on electrically induced contractions in the urethra, and on responses to acetylcholine and electrical field entrehar, and on responses to acetylcholine and electricity in the rabbit. It cannot be excluded that the peptide has a modulatory role in neuronsmission in human urethral muscle citivity in the rabbit. It cannot be excluded that the peptide has a modulatory role in neuronsmission in human urethral muscle citivity in the rabbit.

L12 ANSWER 100 OF 121 ACCESSION NUMBER:

DOCUMENT NUMBER:

AUTTHOR . SOURCE

MEDLINE on STN
85285317 MEDLINE
PubMed ID: 4029260
Pharmacological evidence for selective inhibition
of gastric acid secretion by telenzepine, a new
antimuscarinic drug.
Eltze M; Gonne S; Riedel R; Schlotke B; Schudt C; Simon W A
European journal of pharmacology, (1985 Jun 7) 112 (2)

PUB. COUNTRY

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

ENTRY DATE

HOR: Eltze M, Gonne S, Riedel R; Schlotke B; Schudt C; Simon W A
RCS: European journal of pharmacology, (1985 Jun 7) 112 (2)
211-24.

Journal code: 1254354. ISSN: 0014-2999.

Netherlands
JUNGE: Snglish
ESCHONTY: Netherlands
JUNGE: English
ESCHONTY: Priority Journals
RY MONTH: 198510
RY DATE: Entered STN: 19900320

Last Updated on STN: 19900320

Entered Medline: 19851009
The new antisecretory drug, telenzepine (4,9 dihydro-3-methyl-4-[(4-methyl-1-piperazinyl)acetyl]-10H-thieno-[3/4 - b][1,5]benzdepin 10-one), was investigated for its inhibition of functionally intact
mumcarinic receptors involved in gastric acid secretion in rabbit fundic glands, perfused mouse stomach in vitro, perfused rat stomach in situ, gastric fistula in rats and dogs with a Neidenhain pouch. The effects on these receptors were contrasted with effects on receptors located on smooth muscle and heart, i.e. isolated rat urinary
bladder, stomach and atrium. The results were compared to those values obtained with nonselective antimuscarinic drugs
(N-methylacopolamine, atropine) and the selective M-1 antagonist pirenzepine. Telenzepine was found to be 4-10 times more potent than pirenzepine with respect to depressing both gastric acid secretion and smooth muscle or myocardial responses. Based on log ECS0 and pA2 values, both drugs exhibited a similar selectivity profile differing from the pattern of effects observed with atropine or a second reference compound, solenzepine. As compared with atropine telenzepine exhibited a 5 fold higher relative affinity to muscarinic receptors involved in gastric acid secretion. It was concluded that telenzepine is selective to discriminate between muscarinic receptors mediating gastric acid secretion and affecting muscle contractility and that this finding supports the concept of muscarinic receptor heterogeneity.

L12 ANSWER 102 OF 121 ACCESSION NUMBER: 84 DOCUMENT NUMBER: PU TITLE: Di

ALITUAD

SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH:

ANSWER 102 OF 121 MEDLINE ON STN
ESSION NUMBER: 84076350 MEDLINE
UNENT NUMBER: PubMed ID: 650180
LE: Differences between binding affinities of some antimuscarninic drugs in the parotid gland and those in the wrinary bladder and ileum.

ACR: Nilvebrant L; Sparf B
ACE: Acta pharmacologica et toxicologica, (1983 Oct) 53 (4) 304-13.

JOURNAL Denmark
MENT TYPE: Journal; Article; (JOURNAL ARTICLE)
UNIAGE: English
ESSCMENT: Priority Journals
TY MONTH: 198401
TY MONTH: 198401
TO Sessible differences between the muscarnic receptors in the guinea pig urinary bladder and those in the ileum and the parotid gland were investigated, using a receptor binding technique. The affinities of 19 antimuscarninic drugs were indirectly derived from the ability to inhibit the receptor-specific binding of the radioligand (-)3H-ONB. The Hill coefficients were close to unity which indicated that the drugs were bound to apparently uniform populations of receptors within each tissue. In contrast to traditional muscarnic antagonists, four drugs - namely, oxybutynine, dicyclomine, benzhevol and pirenzepine - bound with a significantly higher affinity in the parotid gland chan in the urinary bladder and ileum coverine. Thus, the present results further support the hypothesis that differences in muscarninic drugs.

112 ANSWER 103 OF 121 MEDLINE on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
DOCUMENT NUMBER:
DOCUMENT NUMBER:
TITLE:

Bubble 15 6308243

TITLE:

Lectrical and mechanical activity recorded from rabbit urinary bladder in response to nerve stimulation.

AUTHOR:

Creed K E; Ishikawa S; Ito Y
JOURNAL Of physiology, (1983 May) 338 149-64.
JOURNAL Of physiology, (1983 May) 338 149-64.
JOURNAL OF SECONDARY:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT Priority JOURNAL ARTICLE)
ENGIAND: United Kingdom
JOURNAL ARTICLE)
ENTRY MONTH:
198309
ENTRY DATE:
English
File SEGMENT:
Priority JOURNAL ARTICLE)
Entered STN: 19900319
Entered Medline: 19830923

AB Responses of the smooth muscle membrane of the rabbit bladder to intramuscular nerve stimulation were investigated by the micro-electrode and double sucrose-gap methods. The cell generated regular spontaneous action potentials. Acetylcholine produced a maintained increase in the frequency and ATP a transient increase. Noradrenaline only increased the frequency at very high concentrations. Application of short current pulses (50 microseconds) produced an initial excitatory junction potential (e.j.p.) with a superimposed spike, followed by a late depolarization. On some occasions, hyperpolarization of the membrane appeared between initial e.j.p. and the late depolarization. All these responses were abolished by tetrodotoxin. The late depolarization was enhanced by pre-treatment with neostigmine and abolished by atropine. This means that the delayed depolarization is due to activation of the membrane appeared between with the e.j.p. and contraction were unaffected by guamethidine, phentolamine, methysergide, mepyramine, quinidine or theophylline. This means that the e.j.p. is not mediated by activation of adrenergic, tryptaminergic, histaminergic or purinergic receptors. ATP reduced the amplitude of the e.j.p. due to depolarization of the membrane and reduction in the membrane resistance. The amplitude of the e.j.p. was gradually reduced by repetitive stimulation of 5-2.0

A

L12 ANSWER 105 OF 121 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR

121 MEDLINE on STN
83174804 MEDLINE
PubMed ID: 6837322
Muscarinic receptor binding in the guinea pig
urinary bladder.
Nilvebrant L; Sparf B
Acta pharmacologica et texicologica, (1983 Jan) 52 (1)
30-8. SOURCE

30-8.
Journal code: 0370572. ISSN: 0001 6683.
Denmark
Journal, Article; (JOURNAL ARTICLE)
English
Priority Journals
198305

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

SEGMENT: Priority Journals
IN MONTH: 198305
IN MONTH: 198305
IN MONTH: 198305
Entered STN: 19900318
Last Updated on STN: 19900318
Entered Medline: 19830505
The muscarinic receptors in the guinea pig urinary
bladder and the longitudinal muscle of the guinea pig ileum were
studied by means of a receptor-binding technique. 1-Quinuclidinyl [phenyl
4-3H]benzilate (-]3H-QNB) was employed as radio-ligand and the separation
of bound from free (-) 3H-QNB was performed by microcentrifugation. Under
conditions of equilibrium (-]3H-QNB was performed by microcentrifugation. Under
conditions of equilibrium (-]3H-QNB was performed by microcentrifugation. Under
conditions of equilibrium (-]3H-QNB was performed by microcentrifugation in
the bladder and ileum, respectively. The binding appeared to
represent a single population of non-interacting binding gites. The
apparent dissociation constants were 2.6 x 10(-10) M in the
bladder and 1.2 x 10(-9) M in the ileum, whereas the KD-values,
estimated by extrapolation to an infinitely low receptor concentration
were 1.1 x 10(-10) M (bladder) and 3.1 x 10(-10) M (leum). The
binding of (-)3H-QNB appears to represent an interaction with
muscarinte receptors, as it was effectively inhibited by
muscarinte antagonists and agonists, but not by a variety of
non-cholinergic drugs.

L12 ANSWER 104 OF 121
ACCESSION NUMBER: 832
DOCUMENT NUMBER: Put
TITLE: CON

AUTHOR

PUB. COUNTRY

DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

L12 ANSWER 106 OF 121 ACCESSION NUMBER: 83

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

ANSWER 106 OF 121 MEDLINE ON STN
ESSION NUMBER: 83147502 MEDLINE

MENT NUMBER: PubMed ID: 7164219

LE: Central cholinergic mechanisms in L DOPA induced hyperactive urinary bladder of the rat.

40R: Sillen U; Rubenson A; H;almas K

CE: Urological research, (1982) 10 (5) 239 43.

JOURNAL COUNTRY: GERMANY, MEST: Germany, Federal Republic of

MENT TYPE: Journal; Article; (JOURNAL ARTICLE)

10AGE: English
10SAGENT: Priority Journals
10SAGENT: Priority Journals
10SAGE: Entered STN: 19900318

10SAGE Entered Medline: 19830407

The involvement of central and peripheral cholinergic structures in the mediation of a centrally induced hyperactive urinary

bladder response to L-2,3-dihydroxyphenylalanine (L-DOPA), after peripheral decarboxylase inhibition, registered by a cystometric procedure, has been analysed pharmacologically in anaesthetised rats. The urinary bladder response to L-DOPA was unchanged after blockade of cholinergic receptors with methylscopolamine, diminished after barropine and totally inhibited after hexamethonium. In addition, activation of muscarinic receptors in the pontine-mesencephalic brain region with oxotremorine after methylscopolamine pretreatment generates a hyperactive urinary bladder response, mediation of which seems to be independent of endogenous catecholamine stores. It is suggested that cholinergic receptors in the pontine-mesencephalic brain region are of importance for regulation of urinary bladder function in the rat.

Puthermore, the bladder hyperactivity urinduced by L-DOPA might be propagated via muscarinic receptors in this brain area, and mediated peripherally via cholinergic receptors in the autonomic ganglia, but in the bladder detrusor via non-cholinergic receptors.

L12 ANSWER 107 OF 121 MEDLINE on STN ACCESSION NUMBER DOCUMENT NUMBER:

121 MEDLINE on SIM
83091092 MEDLINE
FUDMed ID: 6129625
Direct evidence against a role of ATP as the nonadrenergic,
noncholinergic inhibitory neurotransmitter in
guinea pig tenia coli.
Westfall D P; Hogaboom G K; Colby J; O'Donnell J P; Fedan J AUTHOR:

5 T32 GM07039 (NIGMS) CONTRACT NUMBER NBOBBOO

SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

REACT NUMBER: 5 T32 GM07039 (NIGMS)
NB08300
RCE: Proceedings of the National Academy of Sciences of the United States of America, (1982 Nov) 79 (22) 7041-5.
Journal code: 750876. ISSN: 0027-8424.
United States
MENT TYPE: Journal; Article; (JOURNAL ARTICLE)
HORT TYPE: Journal; Article; (JOURNAL ARTICLE)
SEGMENT: Priority Journals
RY MONTH: 198302
RY DATE: Entered Medine: 19830225
Electrical field stimulation of the isolated guinea pig tenia coli in the presence of a muscarinic receptor antagonist (atropine) and an adrenergic neuron blocker (guanethidine) produces relaxation. A large amount of indirect evidence has suggested that the neurotransmitter that is released from these monadrenergic, noncholinergic inhibitory neurons is ATP or a related nucleotide, and the nerves have been termed "purinergic." A photoaffinity analog of ATP, arylazido aminopropionyl ATP, which produces a specific pharmacological antagonism of P2 purinergic receptors in isolated guinea pig vas deferens and urinary bladder, was utilized in the present study to evaluate directly neurotransmitter in tenia coli. By blocking postjunctional P2 receptors, arylazido aminopropionyl ATP produced a pronounced antagonism of relaxations induced by exogenously added ATP. Responses produced by ADP, AMP, and adenosine also were antagonized by arylazido aminopropionyl ATP, but to a lesser extent. Inhibitory responses to isoproterenol were not antagonized. Under these conditions of established, specific P2-receptor blockade of responses to exogenously added ATP, relaxations induced by field stimulation of intrinsic inhibitory nerves in the presence of atropine (1 microM) and guanethidine (1 microM) were not antagonized. Though these results provide no indication of the actual substance involved, they suggest strongly that the nonadrenergic, noncholinergic inhibitory neurotransmitter in the guinea pig tenia coli is not ATP.

ANSWER 109 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

121 MEDLINE on SIN 81200946 MEDLINE PubMed ID: 6112793 Central neurotransmitter mechanisms involved in the control of urinary bladder function. An experimental study in the rat.

AUTHOR SOURCE

Sillen U Scandinavian journal of urology and nephrology. Supplementum, (1980) 58 1-45. Journal code: 0153034. ISSN: 0300-8886.

Journal code. Classis Sweden Journal; Article; (JOURNAL ARTICLE) PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

English Priority Journals 198107 FILE SEGMENT: ENTRY MONTH:

ENTRY DATE:

SIGNAME

English

S SEGMENT: Priority Journals

YMONTH: 198107

XI DATE: Entered STN: 19900316

Last Updated on STN: 19980206

Entered Medline: 19810720

Central neurotransmitter systems have been activated in anaesthetized rats, while the effects on the wrinary bladder have been recorded by cystometric procedures. Stimulation of central catecholamine neurons with the amine precursor L-3.4

dihydroxyphenylalanine (L-DOPA), after peripheral inhibition of the degradating enzyme dopadecarboxylase, resulted in a hyperactive wrinary bladder response. This bladder action seems to be elicited mainly via central dopamine receptors, and mediated via neurons of non-catecholamine type. Activation of central muscarinic receptors with oxotremorine, after pretreatment with methylscopolamine, induced a hyperactive wrinary hladder response, which is suggested to originate in pontine-mesencephalic structures as well. There might in some instances be an interaction with muscarinic receptors in the generation of the bladder response to stimulation of central muscarinic receptors, on the other hand, seems to be independent of intact adrenergic neurons. The peripheral mediation of the bladder response to L-DOPA, at the pontine-mesencephalic brain level. The bladder receptors, but in the bladder detrusor via non-cholinergic as well as non-adrenergic receptors. Activation of central GABA mechanisms with GABA, muscinol and diazepam strongly inhibited the bladder hyperactivity to L-DOPA. This inhibition probably occurred in the pontine-mesencephalic brain area. The results suggest that excitatory dopaminergic and muscarinic receptors, as well as inhibitory gabaergic receptors in the pontine-mesencephalic brain area. The results suggest that excitatory dopaminergic and muscarinic receptors in the pontine-mesencephalic brain area. The results suggest that excitatory dopaminergic and muscarinic receptors in the pontine-mesencephalic brain area.

L12 ANSWER 108 OF 121
ACCESSION NUMBER: 82
DOCUMENT NUMBER: Put
TITLE: Ef:

AUTHOR: SOURCE:

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

RESION NUMBER: 82196037 MEDLINE ON STN

CESSION NUMBER: 92196037 MEDLINE

CLE: 82196037 MEDLINE

CLE: 2821697

CLE: 2821697

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L12 ANSWER 110 OF 121 MEDLINE ON STN
ACCESSION NUMBER: \$1109754 MEDLINE
DOCUMENT NUMBER: \$1109754 MEDLINE
TITLE: Pubmed ID: 7193020
Pharmacology of secoverine, a new spasmolytic agent with specific antimuscarinic properties. Part 1: Antimuscarinic and spasmolytic effects.

AUTHOR: Zwagemakers J M; Claassen V
SOURCE: Arzneimittel-Forschung, (1980) 30 (9) 1517-26.
JOURNAL COUNTRY: DOCUMENT TYPE: JOURNAL ARTICLE)
ENGLAGE: SeffMANY, MSST: Germany, Federal Republic of Journal; Article; (JOURNAL ARTICLE)
English

FUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

COUNTRY: GERMANY, NEST: Germany, Federal Republic of JUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)
SUAGE: English
E SEGMENT: Priority Journals
RY MONTH: 198103
RY DATE: Entered STN: 19900316
Last Updated on STN: 19900316
Entered Medline: 19910317
1-Cyclohexyl-4-[ethyl[p-methoxy-alpha methylphenethyl]amino]-1-butanone hydrochloride (secoverine hydrochloride) is a neurotropic spasmolytic agent with specific antimuscarinic properties. On the basis of the results obtained, the possibility exists that secoverine acts only a sub-group of muscarinic receptors. 1. In vitro experiments the competitive antagonism of secoverine against muscarinomametics was demonstrated on guinea pig ileum, rat jejunum and calif traches smooth muscle. On the basis of the mean difference of the pA2 values and in accordance with the relative activity as determined by 4-point assays, it may be concluded that secoverine is about 0.6 times as active as atropine.
2. In in vive experiments the antimuscarinic activity of secoverine, by both parenteral and intraduodenal routes of administration. It was shown that the action of secoverine was reversible, of quick onset and of long duration. 3. By contrast, secoverine had only marginal effects on the aphincter and ciliary muscle of the eye, almost no effect on cholinergically-induced salivation and lacrimation, gastric acid production, urinary bladder function, gastric emptying or normal peristalsis. 4. The central anticholinergic activity was more in accordance with the activity found in the spasmolytic tests. 5. Apart from the neurotropic action, secoverine has also a good musculotropic activity as was found in in vivo and in vitro experiments. The activity varied from 1.3-13. Itmes that of papaverine in the different organs investigated. The musculotropic activity is not caused by a specific, verapamil-like, calcium antagonism. 6. Secoverine has no nicotinolytic or antihistamanic activity, a moderate antisterotonic activity, an inhibiting effect on the noradrealine uptake mechanism of the vas

09/960.477

L12 ANSWER 111 OF 121
ACCESSION NUMBER: 81
DOCUMENT NUMBER: Pul
TITLE: (Pe MEDLINE on STN 81090827 MEDLINE PubMed ID: 7449999

PubMed ID: 7449998 [Possibility of substance selective action on the m-cholinoreceptors of a specific site]. C vormcohnosti izbiratel'nogo deistviia veshchestv na m-kholinoretseptory opredelennoi lokalizatsii. Kharkevich D A; Skoldinov A P; Samailov D N; Shorr V A Farmakologiia i toksikologiia, (1980 Nov-Dee) 43 (6) AUTHOR:

Journal code: 16920420R. ISSN: 0014-8318.

PUB. COUNTRY: USSR

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE) LANGUAGE: Russian

FILE SEGMENT: Priority Journals ENTRY MONTH: ENTRY DATE: 198103

2 SEGNET: Priority Journals
YM MONTH: 198103
Entered STN: 19900316
Last Updated on STN: 19970203
Entered Medline: 19810327

Effects of a number of agents on the muscarine-sensitive acetylcholine receptors (muscarinic receptors) of different localization were investigated. Experiments on anesthetized cats revealed the following effects of acetylcholine: hypotension, bradycardia, bronchospasm, contractions of the ileum and urinary bladder, hypersalivation. It was shown that the highest sensitivity to bisquaternary ammonium derivatives of diphenyl cyclobutanedicarboxylic (truxillic) acid is exhibited by the muscarinic receptors of the heart. The sensitivity of bronchial muscarinic receptors is slightly lower. The prevailing action on the muscarinic receptors of the heart was also shown by some N-adamanty bisquaternary ammonium compounds. Procaine and the antihistaminics mebhydroline and diphenhydramine eliminated bradycardia induced by acetylcholine without affecting the latter's hypotensive effect. The evidence obtained indicates the heterogeneity of the muscarinic receptors of different localization.

L12 ANSWER 113 OF 121 ACCESSION NUMBER: 80 DOCUMENT NUMBER: Pu 121 MEDLINE on STN 80252482 MEDLINE PubMed ID: 6249955 Cholinergic **inhibition** of **urinary** TITLE:

AUTHOR:

Cholinergic inhibition of urinary acidification by the turtle bladder. Arruda J A; Sabatini S Kidney international, [1980 May] 17 (5) 622-30. Journal code: 0323470. ISSN: 0085-2538. GRMANY, WEST: Germany, Federal Republic of Journal Article; (JOURNAL ARTICLE) COUNTRY:

PUB. COUNTRY: DOCUMENT TYPE:

English

LANGUAGE: FILE SEGMENT: Priority Journals

ENTRY MONTH: ENTRY DATE:

SUAGE: English Priority Journals

8 SEGMENT: Priority Journals

8 Y MONTH: 198010

1 Settle Entered STN: 19900315

Last Updated on STN: 19900315

Last Updated on STN: 19900315

The effect of carbachol on urinary acidification by the turtle bladdar in vitro was studied. Carbachol inhibited urinary acidification in a dose dependent fashion, with half maximal inhibition occurring at 4.5 x 10(-5) M. The effect of carbachol on urinary acidification could be totally prevented by atropine, indicating that the inhibition is mediated through a muscarinic receptor. Carbachol inhibited hydrogen ion secretion by decreasing the active proton conductance and not by altering the proton motive force. Carbachol failed to increase passive loss of hydrogen ion from the mucosa. Carbachol increased calcium uptake by the turtle bladdar; this increase in calcium uptake could be prevented by pretreatment with atropine, pentobarbital, or lanthanum. Pentobarbital or lanthanum blunted the inhibitory effect of carbachol on hydrogen ion secretion. In the presence of low extracellular calcium (0.2 mM), carbachol failed to increase calcium uptake but caused a significant inhibition of hydrogen ion secretion. In the presence of normal calcium concentration, carbachol caused a significant efflux of calcium. These data demonstrate that carbachol inhibits urinary acidification and suggest that the mechanism of this inhibition may be related, at least in part, to changes in cytosolic calcium.

L12 ANSWER 112 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR

CONTRACT NUMBER:

121 MEDLINE ON SIN
81024354 MEDLINE
PubMed ID: 6105562
Inhibition and facilitation in parasympathetic
gauglia of the urinary bladder.
de Groat W C; Bocth A M
NS 07923 (NINDS)
Federation proceedings, (1980 Oct) 39 (12) 2990-6. Ref: 37
Journal code: 0372771. ISSN: 0014-9446.
United States
Journal: Article: (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY:

United States Journal; Article; (JOURNAL ARTICLE) General Review; (REVIEW) English DOCUMENT TYPE:

English Priority Journals 198012 LANGUAGE: FILE SEGMENT:

ENTRY MONTH: ENTRY DATE:

English

E SEGMENT: Priority Journals

RY MONTH: 198012

RY DATE: Entered STN: 19900316

Last Updated on STN: 19970203

Entered Medline: 19801216

Neurons in vesical parasympathetic ganglis receive excitatory and inhibitory inputs from both divisions of the autonomic nervous system. Sacral parasympathetic pathways (cholinergic) provide the major excitatory input to these ganglis via activation of nicotinic receptors. Parasympathetic pathways also activate muscarinic incoming the excitatory and excitatory receptors, which may exert a modulatory influence on transmission. Cholinergic transmission is relatively inefficient when preganglionic nerves are stimulated at low frequencies (clik). However, excitatory postsynaptic potentials (EPSPs) and postganglionic firing markedly increase during repetitive stimulation at frequencies of 1-10 Hz. It is concluded that enhanced transmitter release accounts for the temporal facilitation and that vesical ganglia function as "high pass filters" that amplify the parasympathetic excitatory input to the detrusor muscle during micturition. Transmission in vesical ganglis is also sensitive to advenergic inhibitory and facilitatory synaptic mechanisms elicited by efferent pathways in the hypogastric nerves. The effects of exogenous noreprinephrine indicate that adrenergic inhibitory and previngatic depression of transmitter release although postsynaptic adrenergic hyperpolarizing and depolarizing effects have also been noted. Adrenergic facilitation is mediated by beta receptors as well as unidentified receptors. Norepinephrine also can inhibitor or excite spontaneously active neurons in vesical ganglia. The existence of inhibitory and facilitatory synaptic mechanisms in vesical ganglia provides the basis for a complex ganglionic modulation of the central autonomic outflow to the bladder.

L12 ANSWER 114 OF 121 MEDLINE on STN ACCESSION NUMBER:

79021892 MEDLINE
PubMed ID: 81076
[Effect of acetylcholine on the pituitrin induced osmotic flow of water through the wall of the frog urinary bladder! DOCUMENT NUMBER: TITLE:

AUTHOR:

bladder!.
Vliisnie atsetilkholina na vyzvannyi pituitrinom
osmoticheskii tok vody cherez stenku mochevogo puzyria
liagushki.
Bagrov Ia Iu; Manusova N B
Blulleten' eksperimental'noi biologii i meditsiny, (1978
Sep) 86 (9) 321-4.
Journal code: 0370627. ISSN: 0365 9615.
USSR

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH:

Journal code: 0376627. ISSN: 0365 9615.

Journal code: 0376627. ISSN: 0365 9615.

Journal code: 0376627. ISSN: 0365 9615.

JOURNAL ARTICLE;

JUAGE: Russian

3 SEGMENT: Priority Journals

YMONTH: 197812

YDATE: Entered STN: 19900314

Last Updated on STN: 19900314

Entered Medline: 19781227

The role of intercellular pathways in the ADM dependent water transport was studied on the frog urinary bladder by means of acetylcholine (AC) and other cholinergic compounds. AC (10(-3) M) was found to cause a strong suppression of the pituirin-stimulated water flow. Analogous effect was produced by AC on the commotic flow stimulated by cyclic adenosine monophosphate (cAMP) and theolin. The antipituirin effect was not reproduced either by nicotine, nor by potent M-cholinomimetic agents (methylfurmetide and F-2268), and was not reproduced either by nicotine, nor by potent M-cholinomimetic agents (methylfurmetide and F-2268), and was not reproduced either by microtine, nor by potent M-cholinomimetic agents (methylfurmetide and F-2268), and was not responsible for the antipituitrin effect of AC was completely removed by the anticholinesterase drugs with different mode of action (eserine, proserine, armin, acridine iodmethylate, GD-42) in concentrations of 10 (6)--10 (-3) M. It was concluded that the smooth muscles contraction with the subsequent closure of the intercellular spaces was not responsible for the antipituitrinic action of AC. This effect appears to be connected with cholinesterase activation. A possible role of the phosphoinositides in the water permeability regulation of the urinary bladder wall is discussed.

L12 ANSWER 115 OF 121 MEDLINE on STN ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR

121 MEDLINE on STN
77217117 MEDLINE
PubMed ID: 17940
Uropharmacology: v. choline esters and other
parasympathomimetic drugs.
Finkbeiner A E, Bissada N K; Welch L T
Urology, (1977 Jul) 10 (1) 83-9. Ref: 51
Journal code: 0366151. ISSN: 0090-4295.
United States
Journal; Article; (JOURNAL ARTICLE)
General Review; (REVIEW)
English PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: English Priority Journals FILE SEGMENT:

ENTRY MONTH: ENTRY DATE: 197708

Y MONTH: 197708
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Streed STN: 19900314
Last Updated on STN: 19950206
Entered Medline: 19770825
Various parasympathomimetic drugs are discussed, including the choline esters, bethanechol, carbachol, methacholine chloride, and furtrethonium. Other cholinomimetic agents include muscarine, muscarone, arecholine, and pilocarpine. Anticholinesterase agents inhibit or inactivate acetylcholinesterase enzyme and thus result in a prolonged stimulation of cholinergic receptors by endogenous ACh. Bethanechol is the most widely used parasympathomimetic drug in the United States. Its action is mainly muscarinic with activity largely confined to the urinary bladder and to a lesser degree the gastrointestinal tract. It can be administered only subcutameously or orally, and adequate dosage is necessary for a successful response.

ANSWER 117 OF 121 21 MEDLINE on STN 77113136 MEDLINE 77113136 MEDLINE PubMed ID: 837006

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: Atropine resistance and muscarinic receptors in the rat urinary bladder.

AUTHOR: Carpenter F G

Carpenter F G
British journal of pharmacology, (1977 Jan) 59 (1) 42-9.
Journal code: 7502536. ISSN: 0007 1188.
ENGLAND: United Kingdom
Journal, Article; (JOURNAL ARTICLE) SOURCE:

PUB. COUNTRY:

DOCUMENT TYPE: LANGUAGE:

English Priority Journals 197704 FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

L12 ANSWER 116 OF 121 MEDLINE on STN
ACCESSION NUMBER:
77194773 MEDLINE
DOCLMENT NUMBER:
77194773 MEDLINE
TITLE:
PubMed ID: 866699
THE role of prostaglandins in the excitatory innervation of the rat urinary bladder.
AUTHOR:
Choo L K; Mitchelson F
SOURCE:
Prostaglandins, (1977 May) 13 (5) 917-26.
JOURNAL Odde: 0320271. ISSN: 0090-6980.

United States
DOCUMENT TYPE:
JOURNAL ARTICLE)
LANGUAGE:
English
FILE SEGMENT:
Priority Journals
ENTRY MONTH:
ENTRY MONTH:
Entered Medline: 1977073
AB The possible role of FOs in hyoscine-resistant nerve mediated responses of the rat urinary bladder was investigated. Responses
to electrical stimulation were inhibited by a high
concentration of hyoscine (25 micronmol/1) or by the choline uptake
inhibitors, hemicholinium-3 (500 micrommol/1) and
troxypyrrolidinium (500 micrommol/1). Indomethacin (50 micrommol/1)
produced partial blockade (30%) of responses to electrical stimulation
without markedly affecting responses to acetylcholine and the degree of
blockade was of a similar order in the presence of hyoscine or
troxypyrrolidinium. FOE2 (0.028 -2.8 micrommol/1) produced partial blockade slowly developing increase in tone and spontaneous
activity. Responses to electrical stimulation were at most only slightly
increased in the presence of either PG. However, the PGs always increased
the responses to electrical stimulation after indomethacin, indomethacin
plus hyoscine or indomethacin plus troxypyrrolidinium. Responses to
acetylcholine in the presence of indomethacin were not increased by PGE2.
It is concluded that PGE2 and FZalpha do not function is transmitters
responsible for resistance to anti-musarinic drugs in the
bladder but may exert a modulating effect on nervous transmission.

L12 ANSWER 118 OF 121
ACCESSION NUMBER: 74:
DOCUMENT NUMBER: Pul
TITLE: [Co

121 MEDLINE on STN
74261431 MEDLINE
PubMed ID: 4838789
[Combination of various therapeutic methods in the therapy
of insufficient closure of the bladder in old
women].

AUTHOR: SOURCE:

of insufficient closure of the Damager IN Old women].

Beitrag sur Kombination mehrerer Behandlungsverfahren des insuffizienten Blasenverschlusses bei alten Frauen. Slunsky R
Zentralblatt für Gynakologie, (1974 Feb 22) 96 (8) 225-30.

Journal code: 21820100R. ISSN: 0044-4197.

GERMANY, BAST: German Democratic Republic Journal; Article; (JOURNAL ARTICLE)

German
Priority Journals
197408
Entered STN: 19900310
Last Updated on STN: 19900310
Entered Medline: 19740828

PUB. COUNTRY: DOCUMENT TYPE: LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

L12 ANSWER 119 OF 121 MEDLINE ON STN
ACCESSION NUMBER:
T1TLE:
AUTHOR:
SOURCE:
British journal of urology, (1974 Apr) 46 (2) 187 92.
JOURNAL TYPE:
LANGUAGE:
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L12 ANSWER 121 OF 121 MEDLINE ON STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
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AUTHOR:
AUTHOR:
Chesher G B
AGents and actions, (1970 Mar) 1 (3) 128-32.
Journal code: 0213341. ISSN: 0065-4299.

DUB. COUNTRY:
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L12 ANSWER 120 OF 121 MEDLINE ON STN
ACCESSION NUMBER: 72050602 MEDLINE
DOCUMENT NUMBER: 1 4941704
[Therapy of bladder sphineter incontinence in females]. Therapie der Inkontinenz des Sphinkter vesicae bei der Frau.

AUTHOR: SOURCE:

Theraple der inkomtinenz des Spinistes viellen Prau.
Palmrich A H
Medizinische Klinik, (1971 Oct 15) 66 (42) 1405 9. Ref: 38
JOURNAL code: 0376637. ISSN: 0025-8458.
GERMANY, WEST: Germany, Pederal Republic of
JOURNAL ARTICLE;
General Review; (REVIEW)

PUB. COUNTRY: DOCUMENT TYPE:

LANGUAGE: FILE SEGMENT: ENTRY MONTH: ENTRY DATE:

General Kevlew; (KEVIEW) German Priority Journals 197202 Entered STN: 19900310 Last Updated on STN: 19900310 Entered Medline: 19720202